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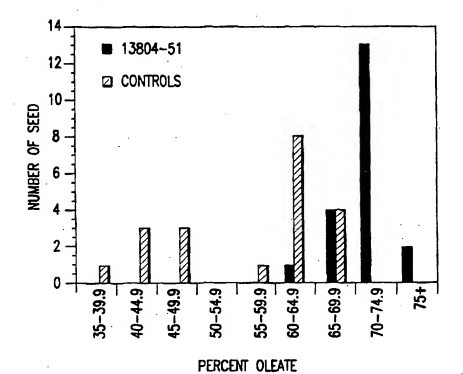
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(57) Abstract

Transformed plants which have increased or decreased linolenic acid content are disclosed. Also disclosed are plants which express a linoleic acid desaturase gene.

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ALTERED LINOLENIC AND LINOLEIC ACID CONTENT IN PLANTS

This is a continuation-in-part of U.S. Serial No. 08/156,551 filed November 22, 1993, which is a continuation of U.S. Serial No. 5 08/014,431, filed on February 5, 1993. The present invention relates to genetically engineered plants. In particular it relates to genetically engineered plants and seeds which have altered linolenic and linoleic acid content compared with naturally occurring plants.

BACKGROUND

10 Many crop species produce seed oils in which the fatty acid composition is not ideally suited to the intended use. The application of conventional breeding methods, coupled in some cases with mutagenesis, has resulted in the production of new varieties of several species with desirable alterations in the fatty acid composition of seed oil. A notable 15 example is the development of low erucic acid varieties of rapeseed (Stefansson 1983). Similar efforts have resulted in the reduction of the level of polyunsaturated 18-carbon fatty acids in soybean (Wilcox and Cavins 1985; Graef et al. 1988), sunflower (Fick 1989), and linseed oils (Green and Marshal 1984).

Most of the genetic variation in seed lipid fatty acid composition appears to involve the presence of an allele of a gene that disrupts normal fatty acid metabolism and leads to an accumulation of intermediate fatty acid products in the seed storage lipids (Downey 1987). However, it seems likely that, because of the inherent limitations of this 25 approach, many other desirable changes in seed oil fatty acid composition may require the directed application of genetic engineering methods.

 α -Linolenic acid (18:3 $^{\Delta9,12,15}$) is an eighteen carbon fatty acid containing three cis double bonds at the 9-10, 12-13 and 15-16 carbons. It is found in the cells of higher plants as a constituent of cell membranes. It is also found in storage organs, such as in seeds. There it is designated oil bodies which are bounded by an electron dense structure that is thought to be a half-unit membrane and dispersed in the cytoplasmic environment of cells. When present as a constituent of cell membranes, linolenic acid is usually esterified to the sn-1 or sn-2 position of the glycerol moiety of a diacyl-glycerolipid. By contrast, when present in oil bodies, linolenic acid is usually esterified to the sn-1, sn-2 or sn-3 position of a triacylglycerolipid (TAG).

Linolenic acid is extensively used in the paint and varnish industry in view of its rapid oxidation. Flax seed is a predominant source of this oil. Soybean seed, on the other hand, does not have sufficient linolenic acid content to be used in this industry. Thus, increasing the linolenic acid content in a plant such as soybean would permit the use of the soybean oil in the paint and varnish industry.

On the other hand, it is undesirable to have significant levels of linolenic acid in cooking oils and foods. Linolenic acid is unstable during cooking and is rapidly oxidized. The oxidized products impart rancidity to the finished product. A rapeseed or soybean oil with reduced linolenic acid, such as containing 2% or less of linolenic acid, would be ideal for use as a cooking oil.

Linolenic acid is also a precursor in the biosynthesis of jasmonic acid, an important plant growth regulator. Linolenic acid is converted to jasmonic acid by introduction of an oxygen to the carbon chain by a lipoxygenase, followed by dehydration, reduction, and several β-oxidations (Vick and Zimmerman, 1984). The activity of jasmonic acid has been measured in terms of induction of pathogen defense responses. By application of free linolenic acid to plants, plant pathogen defenses can also be induced (Farmer and Ryan, 1992).

A model has been proposed to explain the ability of free linolenic acid to exhibit the effects associated with jasmonic acid (Farmer and Ryan, 1992). It is hypothesized that all of the enzymatic activities which are required for the conversion of linolenic acid to jasmonic acid are constitutively present in the cell and the rate limiting step in the production of jasmonic acid is the availability of free linolenic acid. A likely route for the production of the free linolenic acid is by the activity of a lipase in the plasma membrane.

It has been observed that exogenous jasmonic acid can more powerfully activate defense responses than can wounding. This suggests that wounds cannot generate enough free linolenic acid to support high level production of jasmonic acid. The activity of the lipase or the availability of appropriate substrate for the lipase may be rate limiting upon wounding. Thus, increasing the linolenic acid content of plasma membrane may positively influence "signal transduction" in plants and result in better protection against environment and pathogen stress.

Linolenic acid, as well as oleic and linoleic acids are also important constituents, as well as precursors of volatile carbonyl compounds, whic contribute to the aroma of both fresh and cooked foods.

The major fatty acids of tomato fruit pericarp are oleic, linoleic and linolenic acids. As the fruit ripens, the levels of the latter two fatty acids decline resulting in the production of a number of 4-6 carbon containing aldehydees and ketones. One particular metabolite, cis-3-hexanol, has been shown to be present in higher levels in vine-ripened tomatoes compared to supermarket tomatoes or tomatoes stored in refrigerators. It is likely, therefore, that the "aroma" of fresh fruits and vegetables can be "modulated" by regulation of the content of linolenic and linoleic acids, important substrates for the enzyme lipoxygenase and subsequently the

hydroperoxide cleaving enzyme, which generates the volatile "aroma" compounds.

From the above, it is clear that the ability to vary the content of linolenic acid in plants would be desirable. However, to achieve this result it is necessary to determine what controls the product of linolenic acid in plants.

A large body of experimental evidence derived from radiochemical tracer studies has indicated that α-linolenic acid is synthesized by the desaturation of linoleic acid (18:2^{Δ9,12}) (reviewed in Harwood 1988;). However, the actual substrate for desaturation is not known.

In vivo and in vitro labelling studies suggest that there are possibly two distinct pathways for the synthesis of linolenic acid (Browse and Somerville, 1991). One possible pathway is thought to be located in the endoplasmic reticulum where linoleic acid esterified to the sn-2 position of phosphatidylcholine is a substrate for desaturation. However, the available evidence does not exclude the possibility that linoleic acid esterified to other lipids may also be a substrate.

A second possible pathway of linoleic acid desaturation is located in the plastid where the available evidence suggests that linoleic acid esterified to monogalactosyldiacylglycerol and, possibly, other plastid lipids is the substrate for desaturation.

Relatively little direct information is available concerning the enzymes involved in linoleic acid desaturation. Low levels of enzyme activity have been detected in microsomal membrane preparations from developing linseed (Linum ussitatum) (Browse and Slack, 1981) and, more recently, in preparations of gently lysed chloroplasts (Schmidt and Heinz, 1990a,b). The general features of the enzyme may be inferred from information available about other enzymes of this class.

The most thoroughly characterized desaturase is the stearoyl-Coenzyme A (CoA) desaturase from vertebrate liver (reviewed by Holloway, 1983). This enzyme has been shown to be an integral membrane protein which contains non-heme iron. The desaturase reaction requires fatty acyl-CoA, molecular oxygen and reduced cytochrome b5, another membrane protein. In vivo, the reduced cytochrome b5 is produced by the transfer of reducing equivalents from NADH via the activity of cytochrome b5 reductase, a flavin containing membrane protein.

The most thoroughly characterized desaturase from plants is the stearoyl-ACP desaturase (McKeon and Stumpf, 1982; Shanklin and Somerville, 1991). This enzyme also requires molecular oxygen and a high potential reductant. However, in contrast to the animal enzyme, this desaturase is a soluble plastid protein which preferentially acts on a fatty acid esterified to acyl carrier protein (ACP) rather than CoA. This enzyme also differs from the animal enzyme by utilizing reduced ferredoxin as an intermediate electron donor.

Other plant desaturases appear to be membrane proteins. The microsomal $\Delta 12$ oleate desaturase from several plant species has been assayed in membrane preparations from several plants (Harwood, 1988).

20 As with the stearoyl-CoA desaturase from animals, this enzyme requires molecular oxygen and reduced cytochrome b5 as an electron donor (Kearns et al., 1991). However, it appears that oleate esterified to a phospholipid is the substrate rather than a CoA ester.

With regard to the activity responsible for the making of linolenic acid, little was known as to its source or origin. However, evidence that the amount of linolenic acid is related to the amount of linoleic acid desaturase activity has been obtained by analysis of the properties of the fad3 mutant of Arabidopsis thaliana (Lemieux et al. 1990). This mutant is deficient in linolenic acid in the storage oils of its seed lipids and in the

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membrane lipids of different tissues to varying degrees. The mutant also had an increase in the amount of linoleic acid. This can be interpreted as evidence that the mutant is defective in the activity of a desaturase which converts linoleic acid to linolenic acid.

There is further evidence to suggest that the activity of this desaturase could be rate limiting for linolenic acid synthesis under normal circumstances. This was discovered by measuring the effects on fatty acid composition in heterozygous plants (i.e., fad3+/fad-) formed by crossing the wild type with the fad3 mutant. In these F1 plants, which have one copy of 10 the normal fad3 gene product instead of the two normally found in the wild type, the amount of linolenic acid was almost exactly intermediate between that found in either parent. This suggests that the amount of linolenic acid is proportional to the amount of functional fad3 gene product (Lemieux et al., 1990).

These results do not shed any light, however, on the nature of the fad3 gene product or whether the observed effects in mutants are related to either a decrease in quantitiy of desaturase protein or desaturase activity due to a defective protein.

Moreover, nothing is known with any degree of certainty 20 about the linoleic acid desaturase from plant microsomes. As noted above, very little is known about the microsomal desaturases except that they probably utilize reduced cytochrome b5 as intermediate electron donor and probably utilize lipids rather than CoA or ACP esters as substrates.

Moreover, as in many other aspects of plant biology, the lack 25 of specific information about the biochemistry and regulation of lipid metabolism makes it difficult to predict how the introduction of one or a few genes might usefully alter seed lipid synthesis.

An additional problem arises from the fact that many of the key enzymes of lipid metabolism are membrane-bound and present in low

quantities. Thus, attempts to solubilize and purify them from plant sources have not been successful.

SUMMARY OF THE INVENTION

The present invention provides structural coding sequences encoding linoleic acid desaturase activity which can be used to alter the linoleic and linolenic acid compositions of plants or to isolate other plant linoleic acid desaturases. The present invention further provides a plant capable of expressing a structural coding sequence to control the level of linolenic acid or linoleic acid or both in the plant. The present invention further provides a method for controlling the levels of linoleic and linolenic acid in plants. It is also demonstrated by the present invention that the linoleic acid desaturase enzyme activity in plant cells and tissues is a controlling step in linolenic acid biosynthesis.

The present invention further relates to the engineering of two
advantageous traits into plants: increased and decreased α-linolenic acid
content in the structural lipids or storage oils of various crop plants.

In accomplishing the foregoing, there is provided, in accordance with one aspect of the present invention, a genetically transformed plant which has an elevated linolenic acid content comprising a recombinant, double-stranded DNA molecule comprising

- (i) a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;
- (ii) a structural coding sequence that causes the production of an RNA sequence that encodes a linoleic acid desaturase activity; and
- (iii) a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence.

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In accordance with another aspect of the present invention, there is provided a genetically transformed plant which has a reduced linolenic acid content, comprising a recombinant, double-stranded DNA molecule comprising

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- a promoter that functions in plant cells to cause (i) the production of an RNA sequence, said promoter operably linked to;
- a DNA sequence that causes the production of an RNA sequence that is in antisense orientation to at least a portion of a gene that encodes a linoleic acid desaturase activity in said plant; and
- a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence.

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There has also been provided, in accordance with another aspect of the present invention a method of producing a genetically transformed plant which has an elevated or reduced linolenic acid content. There has also been provided, in accordance with another aspect of the present invention a recombinant, double-stranded DNA molecule and plant cells 20 containing a recombinant, double-stranded DNA molecule.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows the genetic map of the region of chromosome 2 of Arabidopsis thaliana where a linoleic acid desaturase gene is located and the identity of the yeast artificial chromosomes which carry this region of 25 the genome.

Figure 2 shows the structure of plasmid pBNDES3 which was obtained by inserting an EcoRI fragment containing the B. napus linoleic acid desaturase cDNA (fad3) into pBLUESCRIPT.

Figure 3 shows the nucleotide sequence (SEQ ID NO:1) and deduced amino acid sequence (SEQ ID NO:2) for the linoleic acid desaturase cDNA (fad3) from *B. napus*.

Figure 4 shows a comparison of the deduced amino acid sequence of one linoleic acid desaturase cDNA (fad3) from B. napus and the desA gene from Synechocystis. Identical residues are indicated by a solid box. Conservative substitutions are indicated by a stippled box.

Figure 5 shows the binary Ti plasmid vector pBI121.

Figure 6 shows the binary Ti plasmid pTiDES3 which was 10 constructed by insertion of a linoleic acid desaturase cDNA (fad3) into pBI121.

Figure 7 shows the map of the plant transformation vector pMON13804.

Figure 8 shows the map of the plant transformation vector pMON13805.

Figure 9 shows the oil content of control and transformed canola seed in accordance with the present invention.

Figure 10 shows the nucleotide sequence (SEQ ID NO:9) for the linoleic acid desaturase cDNA (fadD) from *Arabidopsis*.

Figure 11 shows the deduced amino acid sequence (SEQ ID NO:10) for the linoleic acid desaturase cDNA (fadD) from *Arabidopsis*.

Figure 12 shows the nucleotide sequence (SEQ ID NO:11) for the linoleic acid desaturase cDNA (fadE) from *Arabidopsis*.

Figure 13 shows the deduced amino acid sequence (SEQ ID NO:12) for the linoleic acid desaturase cDNA (fadE) from Arabidopsis.

DETAILED DESCRIPTION OF THE INVENTION

A genetically transformed plant of the present invention which has an altered linolenic or linoleic acid content can be obtained by expressing the double-stranded DNA molecules described in this application.

The expression of a double-stranded DNA involves transcription of messenger RNA (mRNA) from one strand of the DNA by RNA 5 polymerase enzyme, and the subsequent processing of the mRNA primary transcript inside the nucleus. This processing involves a 3' non-translated region which adds polyadenylate nucleotides to the 3' end of the RNA.

Promoters

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Transcription of DNA into mRNA is regulated by a region of 10 DNA usually referred to as the "promoter." The promoter region contains a sequence of bases that signals RNA polymerase to associate with the DNA, and to initiate the transcription of mRNA using one of the DNA strands as a template to make a corresponding complementary strand of RNA.

Any promoter which is known or is found to cause transcription of RNA in plant cells can be used in the present invention. Promoters which are useful in the present invention include any promoter that functions in a plant cell to cause the production of a RNA sequence. A number of promoters which are active in plant cells and are capable of 20 producing a RNA sequence have been described in the literature. These include the nopaline synthase (NOS) and octopine synthase (OCS) promoters (which are carried on tumor-inducing plasmids of Agrobacterium tumefaciens), the caulimovirus promoters such as the cauliflower mosaic virus (CaMV) 19S and 35S and the figwort mosaic virus 35S-promoters, 25 the light-inducible promoter from the small subunit of ribulose-1,5-bisphosphate carboxylase (ssRUBISCO, a very abundant plant polypeptide), and the chlorophyll a/b binding protein gene promoter, etc. All of these promoters have been used to create various types of DNA constructs

which have been expressed in plants; see, e.g., PCT publication WO 84/02913 (Rogers et al., Monsanto).

Promoters may be obtained from a variety of sources such as plants and plant viruses. Promoters can be used in the form that they exist as isolated from plant genes such as ssRUBISCO genes, or can be modified to improve their effectiveness, such as with the enhanced CaMV35S promoter.

Those skilled in the art will recognize that the amount of linoleic acid desaturase needed to induce the desired alteration in linolenic acid content may vary with the type of plant. It is also possible that extremes in linoleic acid desaturase activity may be deleterious to the plant. Therefore, in a preferred embodiment, promoter function should be optimized by selecting a promoter with the desired tissue expression capabilities and approximate promoter strength and selecting a transformant which produces the desired linoleic acid desaturase activity in the target tissues.

This selection approach from the pool of transformants is routinely employed in expression of heterologous structural genes in plants since there is variation between transformants containing the same heterologous gene due to the site of gene insertion within the plant genome. (Commonly referred to as "position effect").

In a preferred embodiment, the promoters utilized in the doublestranded DNA molecules should have relatively high expression in tissues where the increased or decreased linolenic acid content is desired, such as 25 the seeds of the plant. In Canola, a particularly preferred promoter in this regard is the seed specific promoter described herein in greater detail in the accompanying examples.

In another preferred embodiment, the promoter used in the expression of the double-stranded DNA molecules of the present invention

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can be a constitutive promoter, expressing the DNA molecule in all or most of the tissues of the plant. However, the promoter selected for this embodiments should not cause expression at levels which are detrimental to plant health, growth and development.

B-conglycinin (also known as the 7S protein) is one of the major storage proteins in soybean (Glycine max) (Meinke et al., 1981). The 7S (β conglycin) a'-subunit promoter, used in one aspect of this study to express the linoleic acid desaturase gene, has been shown to be both highly active and seed-specific (Doyle et al., 1986 and Beachy et al., 1985). The \(\beta\)-subunit 10 of B-conglycinin has been expressed, using its endogenous promoter, in the seeds of transgenic petunia and tobacco, showing that the promoter functions in a seed-specific manner in other plants (Bray et al., 1987). The promoter for \(\beta\)-conglycinin could be used to in accordance with the present invention. If used, this promoter could express the DNA molecule 15 specifically in seeds, which could lead to an alteration in the linolenic acid content of the seeds.

In addition, the endogenous plant linoleic acid desaturase promoters can be used in the present invention. These promoters should be useful in expressing a linoleic acid desaturase gene in specific tissues, such 20 as leaves, seeds or fruits. A number of other promoters with seed-specific or seed-enhanced expression are known and are likely to be expressed in seeds, which are oil accumulating cells. For illustration, the napin promoter and the acyl carrier protein promoters have been utilized in the modification of seed oil by antisense expression (Knutson et al., 1992).

The linolenic acid content of root tissue can be increased by expressing a linoleic acid desaturase gene behind a promoter which is expressed in roots. The promoter from the acid chitinase gene (Samac et al., 1990) is known to function in root tissue and could be used to express the linoleic acid desaturase in root tissue. Expression in root tissue could also be accomplished by utilizing the root specific subdomains of the CaMV35S promoter that have been identified. (Benfey et al., 1989). The linolenic acid content of leaf tissue can be increased by expressing the linoleic acid desaturase gene using a leaf active promoter such as ssRUBISCO promoter or chlorophyll a/b binding protein gene promoter.

The linolenic acid content of fruits can be increased by expressing a linolenic acid desaturase gene behind a promoter which is functional in fruits. Such promoters could be either expressed at all developmental stages of the fruit or restricted to specific stages, particularly fruit ripening.

The RNA produced by a DNA construct of the present invention can also contain a 5' non-translated leader sequence. This sequence can be derived from the promoter selected to express the gene, and can be specifically modified so as to increase translation of the mRNA. The 5' non-translated regions can also be obtained from viral RNAs, from suitable eukaryotic genes, or from a synthetic gene sequence. The present invention is not limited to constructs, as presented in the following examples, wherein the non-translated region is derived from the 5' non-translated sequence that accompanies the promoter sequence. Rather, the non-translated leader sequence can be derived from an unrelated promoter or coding sequence as discussed above.

Linoleic Acid Desaturase Structural Coding Sequences

The structural coding sequence that causes the production of an RNA sequence that encodes a linoleic acid desaturase activity can be the sequences disclosed in the present application, or any sequence that can be obtained using the sequences disclosed in the present application, or any sequence that can be isolated using the method disclosed in the present application.

The structural coding sequence can also be a part of or from the structural coding sequences disclosed in the present invention. It is possible that the active part of the linoleic acid desaturase is formed using only part of the structural coding sequences disclosed in the present application.

The structural coding sequences can be obtained from a variety of sources, such as algae, bacteria or plants. Preferably, structural coding sequences obtained from plants are used in accordance with the present invention.

Since virtually nothing was known about the properties of the linoleic acid desaturase structural coding sequence prior to the present invention, the method used in the present invention to isolate the structural coding sequence was based on the concept of map based cloning. The essential concept in map based cloning is to use information about the genetic map position of a structural coding sequence to isolate the region of the chromosome surrounding the structural coding sequence, and then to use the isolated DNA to complement a mutation in the structural coding sequence. This strategy has never previously been reported in the isolation of any plant gene.

In order to implement map based cloning of the linoleic acid desaturase, mutants of Arabidopsis thaliana (L.) deficient in linoleic acid desaturase activity were isolated by screening randomly chosen individuals from mutagenized populations of plants for individual plants with altered leaf or seed fatty acid composition. (Browse et al. 1985; Lemieux et al. 1990). By screening thousands of plants for altered fatty acid composition, mutants with decreased amounts of linolenic acid and increased amounts of linoleic acid in leaf and seed lipids were isolated. Physiological and genetic analyses of these mutants indicated that they fell into three complementation groups designated fad3, fadD and fadE.

The fad3 mutants had very reduced levels of linolenic acid in seeds and roots but had almost normal levels of linolenic acid in leaves. This effect was interpreted as evidence that the fad3 locus encoded a microsomal desaturase which was responsible for desaturation of linoleic acid to linolenic acid on lipids made by the pathway of lipid biosynthesis in the endoplasmic reticulum, designated the "eukaryotic pathway" (Lemieux et al. 1990). This pathway is mostly responsible for the synthesis of lipids in non-green tissues such as seeds and roots, but plays a secondary role in leaves and other green tissues. Thus, a mutation in the fad3 gene would not be expected to have a major effect on the desaturation of leaf lipids.

In contrast to the fad3 mutant, the fadD mutant had almost normal fatty acid composition of roots and seeds, but had a strong reduction in the amount of linolenic acid in leaf lipids, and a corresponding increase in the amount of linoleic acid. (Browse et al., 1986). Thus, this mutant had the properties expected of a mutant deficient in a linoleic acid desaturase from the prokaryotic pathway which is primarily responsible for the synthesis of lipids in green tissues.

An unusual property of the fadD mutants was that they were very deficient in linoleic acid content when grown at temperatures above about 22 °C but had almost normal fatty acid composition when grown at temperatures below about 18 °C (McCourt et al., 1987). Since it was very unlikely that several independently isolated mutations would all give rise to a temperature conditional phenotype, it was concluded that a second desaturase must be partially responsible for desaturating linoleic acid to linolenic acid in green tissues. Therefore, the fadD mutant was remutagenized with ethylmethane sulfonate, self-fertilized to produce a segregating population of mutagenized plants (designated the M2 generation), and this population was screened for a mutant which was deficient in linolenic acid in green tissues at low temperatures. A mutant

with this property was isolated and the mutation responsible for this effect was designated the fadE locus (Somerville and Browse, unpublished).

Isolation of the Linoleic Acid Desaturase Gene from Canola

The following example was used to isolate the structural coding sequence from the fad3 region. The method described herein could equally have been used to isolate either the fadD or fadE region.

In order to approximately locate the fad3 mutation of the genetic map of Arabidopsis, a sexual cross was made between the fad3 mutant line BL1 and the multiply marked mutant line W1 (Hugly et al., 1991). The F1 hybrids from this cross were permitted to self-fertilize and the resulting F2 plants were scored for both the segregating genetic markers and the altered fatty acid composition. The results of this analysis indicated that the fad3 mutation was located on chromosome 2 near the marker erecta. In order to obtain a more accurate map position by RFLP mapping, a second sexual cross was made between the fad3 mutant line BL1 and the Niederzenz race of Arabidopsis. The F1 progeny were permitted to self-fertilize to produce the F2 generation. 137 F2 plants were grown during 3 weeks at 22° C (100 µE/m²/s) in order to produce fully expanded rosettes, and a few leaves (representing a total weight of 0.2-0.5 g per plant) were harvested from each plant in order to prepare DNA from them.

The leaves were frozen in liquid nitrogen, and ground in dry ice, using a mortar and a pestle. For each sample, the frozen powder was transferred to a microfuge tube and an equal amount of 2 X CTAB buffer (2% cetyltrimethyl ammonium bromide (CTAB), 100 mM Tris-HCl pH 8, 20 mM EDTA, 1.4 M NaCl, 1% polyvinylpolypyrrolidone (PVP) 40,000) was added. The tubes were left at room temperature for 5 min to allow the powder to thaw. The homogenate was extracted once with a mixture of chloroform-isoamyl alcohol (24:1, v/v), and 1/10 vol of 10 X CTAB (10 % CTAB, 0.7 M NaCl) buffer was added to the aqueous phase, which was then

reextracted with an equal volume of chloroform isoamyl alcohol (24:1, v/v). The aqueous phase was transferred to a fresh microfuge tube and 1.5 vol of CTAB precipitation buffer (1% CTAB, 50 mM Tris-HCl pH 8, 10 mM EDTA) was added. The DNA was allowed to precipitate for 12 hr at 4 degrees, and collected by centrifugation (5 min at 10 000g). The DNA was resuspended in 100 µl of 10 mM Tris-HCl pH 7.5, 1 mM EDTA, 1 M NaCl, and 100 µg/ml RNase A and incubated at 50°C for 30 min. The DNA was precipitated by adding 2.2 vol of ethanol and incubating on ice for 20 min. The DNA was collected by centrifugation and the pellet was washed once with 1 ml of 70% ethanol, dried under vacuum for 3 min and resuspended in 10 µl of distilled water. The DNA was stored at -20°C until use.

The 137 plants were grown to maturity and their seeds were collected individually. The fatty acid composition of 10 individual seeds from each of the F2 plants was measured as described by Browse et al (1986) in order to score the fad3 phenotype of each plant. Each seed was incubated in 1 ml of 1N HCl in methanol for 1h at 80°C. The tubes were cooled to room temperature and 1 ml of 0.9 % NaCl plus 0.3 ml of hexane were added. The tubes were agitated by vortexing and the phases separated by centrifugation (300xg for 5 min). The hexane phase was saved, evaporated under a stream of nitrogen, and the fatty acid methyl esters were dissolved in 50 µl hexane. An aliquot (2 µl) was injected onto the gas chromatograph and the fatty acid methyl esters separated and quantitated by flame ionization as described (Browse et al., 1986).

The DNA samples (1 µg) were then cut with the appropriate restriction enzyme (EcoR1 for the marker # 220, Bgl2 for the marker ASA2) using a concentration of 1XKGB buffer (Sambrook et al, 1989), 5 units of the restriction endonuclease and 100 µg/ml BSA. The volume of each sample was 10 µl and the incubation was performed at 37 °C for 4 h. The fragments were resolved by agarose gel electrophoresis (0.8 % agarose

in 1X TAE buffer; Sambrock et al., 1989) and transferred to nylon filters (hybond N+), using the alkaline transfer method as described by the manufacturer. The nylon filters were probed (according to Church and Gilbert, 1984) with radioactively labelled fragments of DNA (Sambrock et al., 1989) corresponding to known RFLP markers which had previously been mapped in the approximate vicinity of the fad3 locus on chromosome 2. The RFLP markers 220 (Chang et al 1988) and ASA2 were found to map close to the fad3 locus. Analysis of the pattern of recombinants (Table 1) indicated that both ASA2 and 220 were located on the same side of the fad3 locus at distances of 0.4 and 2.2 centimorgans (cM), respectively.

		Ta		
	# of plants	<u>220</u>	ASA2	fad3
	67	H	H	+/-
15	30	L	L	-/-
	34	N	N	+/+
	3	H	N	+/+
	1	L	H	+/-
,	1	N	H	+/-
20	1	H	H	-/-

Table 1 shows the genotype of the F2 plants used for mapping the fad 3 locus. L is for Landsberg (background of the fad 3 mutant), N is for Niederzenz, H for heterozygous. A total of 137 F2 plants were analyzed.

The number of recombinant plants between fad3 and 220 or ASA2 was 6 and 1 respectively.

In order to isolate the region of the chromosome containing the fad3 locus, the RFLP markers 220 and ASA2 were used as hybridization probes to screen several yeast artificial chromosome (YAC) libraries. (Grill

and Somerville, 1991; Ward and Jen, 1990). The YAC filters were prepared according to Grill and Somerville (1991). The library was replicated onto nylon filters disposed on petri dishes of SC — (synthetic complete medium minus tryptophan and uracil; Sherman et al., 1986). The cells were allowed to grow for 12 h at 30°C, and the filters were transferred for 15 min on a Whatman 3MM paper saturated with 1 M sorbitol, 50 mM DTT, 50 mM EDTA (pH 8).

The cell wall of the cells was then digested with lyticase, by incubating the filters on a Whatman paper saturated with 1M sorbitol, 50 mM EDTA and 2 mg/ml lyticase (Sigma Co., St. Louis, MO) for 12 h at 30°C. The filters were then transferred on a Whatman 3MM paper saturated with 0.5 M NaOH, 1.5 M NaCl for 15 min, neutralized with 0.5 M Tris-HCl pH 8 for 15 min and quickly rinsed in 2XSSC (SSC is 10mM sodium citrate, 150mM NaCl, pH 7). The filters were allowed to dry, and were transferred to a vacuum oven at 80°C for 1 h. They were subsequently hybridized according to Church and Gilbert (1984), with probes labelled with 32P according to Sambrook et al. (1989).

The DNA of RFLP probe 220 was prepared from 100 ml of liquid culture lysate using the lambdasorb procedure (Promega Corp., Madison, 20 WI); the cDNA encoding ASA2 was excised from the original plasmid (pKN140C; obtained from Dr. G. Fink, Whitehead Institute, Cambridge, MA) with Hind3 and cloned into the Hind3 site of pBLUESCRIPT. The plasmid DNA was then purified by Cesium chloride gradients according to Sambrook et al (1989), digested with Hind3 and the DNA insert was gel purified twice by electroelution according to Sambrook et al (1989).

In order to probe the libraries, the whole DNA from RFLP220 was used as a hybridization probe. By contrast, only the DNA insert of ASA2 was used as a probe. The RFLP probe 220 hybridized to YAC

EG4E8 and EG9D12. The probe ASA2 hybridized to YACs EW15G1, EW15B4 and EW7D11.

In order to determine if these YACs contained all of the DNA between RFLP220 and ASA2, small regions of DNA from the ends of the inserts in EG4E8 and EW15G1 were prepared by inverse PCR (Grill and Somerville, 1991). For that purpose, DNA was prepared from the appropriate YAC clones. The clones (single colonies) were grown to saturation in SC-- liquid cultures, and 1 ml of these cultures was used to inoculate 40 ml liquid cultures (in SC-- medium) that were allowed to grow for 16 h at 30°C. The cells were collected by centrifugation, washed once in 1 M sorbitol, 50 mM EDTA, resuspended in 200 μl of 1 M sorbitol, 50 mM EDTA, 100 mM sodium citrate pH 5.8, 2 mM β-mercaptoethanol and 2 mg/ml lyticase, and incubated 2 h at 30 °C.

Next, 350 µl of 2XCTAB buffer was added and the DNA was purified as described above. DNA (5 µg) of each clone was digested separately with HincII, AluI, EcoRV and RsaI (in 1XKGB buffer, at 37 °C for 4 h; final volume: 50 µl). The reactions were stopped by heating at 65 °C for 15 min, extracted once with one volume of phenol saturated with TE pH 8, followed by an extraction with 1 volume of chloroform - isoamyl alcohol mixture (24:1, vol/vol). The DNA was recovered by ethanol precipitation and resuspended in sterile distilled water. The ligation reactions were performed using 300 ng of DNA in a final volume of 50 µl. The reactions were carried out in 50 mM Tris-HCl pH 7.4, 10 mM MgCl2, 1 mM DTT,1.2 mM ATP with 1 U of ligase, for 2 h at 20 °C, and stopped by heating at 68 °C for 30 min.

The PCR reactions were carried out as follows: The buffers used were the ones indicated by the suppliers except for the Perkin Elmer enzyme for which the reaction was supplemented with an additional 1.4 mM MgCl₂ (final concentration 2.9 mM Mg). The dNTP final concentration

was 125 µM when the Perkin Elmer enzyme was used and 200 µM with the Taq polymerases from other sources. In all cases, 100 ng of each oligonucleotide was used. The final volume was 100 µl. When no product was obtained, the reactions were carried out again in the same conditions except that formamide was added to a final concentration of 3 %.

The left end was amplified from the ligation products of the EcoRV and RsaI digests, using the oligonucleotides EG1 (GGCGATGCTGTCGGAATGGACGATA) (SEQ. ID NO. 3) and EG2 (CTTGGAGCCACTATCGACTACGCGATC) (SEQ. ID NO. 4).

The right end of the clones obtained from the EG library was amplified from the ligation products of the AluI and HincII digests, using the oligonucleotides EG3 (CCGATCTCAAGATTACGGAAT) (SEQ. ID NO. 5) and EG4 (TTCCTAATGCAGGAGTCGCATAAG) (SEQ. ID NO. 6).

The right end of the clones obtained from the EW YAC library was amplified using the oligonucleotides H1 (AGGAGTCGCATAAGGGAG) (SEQ. ID NO. 7) and H2 (GGGAAGTGAATGGAGAC) (SEQ. ID NO. 8), using the same cycle conditions as above, except that the annealing temperature was reduced to 50 °C.

After the reactions were completed, 5µl of each mixture were electrophoresed on an agarose gel to separate the amplification product from primers. The slice of agarose that contained the amplified band was excised from the gel and melted in 1 ml of distilled water. Large amounts of product could then be produced, by reamplification of 5 µl of the melted slice. The PCR products were then purified by electroelution or by using GeneClean (Bio101) and used as hybridization probes to probe filters containing the isolated YAC DNA restricted by several enzymes. The probe made from the right end of EW15G1 hybridized to EG4E8 and similarly, a probe from the right end of EG4E8 hybridized to EW15G1.

Thus, it was concluded that the YACs EG4E8 and EW15G1 contained all of the DNA in the region of the chromosome between RFLP220 and ASA2.

The size of the YAC clones was estimated by field inversion electrophoresis (CHEF, Vollrath and Davis, 1987). High molecular weight 5 DNA was prepared as follows: the yeast cells which contained the YAC clones were grown and treated with lyticase as for preparing DNA as described above. The spheroplasts were then resuspended in an equal volume of 1M sorbitol, 50 mM EDTA, 1% low melt agarose at 37°C. The mixture was poured in a mould (Biorad) which was set on ice to allow the agarose to harden.

The resulting plugs were incubated for 12 h in 0.5 M EDTA pH 9, 1% lauryl sarcosine 1 mg/ml Proteinase K at 50°C. The plugs were subsequently washed twice in 50 mM EDTA and stored at 4°C until use. The CHEF gel was run in 1XTBE for 16 h at 200 V, with a switching 15 interval of 20 s; the temperature of the buffer was maintained at 14 °C during the run. The sizes of the YACs were determined by comparison with a lambda ladder and the yeast chromosomes, and were as follows: EG4E8, 90 kb; EG9D12, 190 kb; EW15G1, 90 kb; EW15B4, 70 kb, EW7D11, 125 kb. These sizes permitted us to roughly determine a correspondence 20 between physical and genetic distances: the distance that separates 220 from ASA2 cannot exceed 180 kb, the sum of the size of the 2 YACs EG4E8 and EW15G1. Since the corresponding genetic distance is 1.7 cM, one can roughly estimate that, in this particular cross and in this particular region of the genome, the value of 1 cM is close to 100kb. Thus, since the 25 fad3 gene maps only 0.4 cM away from ASA2, the corresponding physical distance should be close to 40 kb. We then concluded that fad3 was probably located on the YAC EW7D11, which is the largest YAC hybridizing with ASA2. See Figure 1.

In order to test the possibility that the YAC EW7D11 carried the fad3 gene, the YAC was used to probe a cDNA library made from developing seeds of Canola (Brassica napus L.). Even though the YAC was isolated from Arabidopsis, the fact that Arabidopsis and B. napus are both 5 members of the family Cruciferae led us to predict that the homologous genes from these two species would be sufficiently identical at the nucleotide sequence level so that the Arabidopsis gene would hybridize to the B. napus gene. We also assumed that, because it catalyzes a chemically similar reaction to the stearoyl-ACP desaturase, it would be 10 expressed at similar moderately high levels in developing seeds (Shanklin and Somerville, 1991). Since EW7D11 contained only about 0.2% of the total genome, we expected it to contain only about 2 moderately abundantly expressed genes (i.e., genes in which the mRNA is between 0.1 and 0.01% of total mRNA).

DNA of YAC EW7D11 was isolated as follows: high molecular weight DNA was prepared from the yeast cells that contained the YAC EW7D11 as described above, and several preparative low-melt agarose CHEF gels were run in 1XTBC buffer (same as TBE except that CDTA was substituted for EDTA). The slices that contained the YAC were excised 20 from the gels and pooled. Three slices were melted at 65°C and extracted with an equal volume of phenol saturated with TE. The aqueous phase was saved and reduced to 0.5 ml by repeated extractions with isobutyl alcohol. The remaining agarose was removed by several phenol extractions, followed by two chloroform-isoamyl alcohol extractions. The DNA was precipitated 25 by adding 2 μg of linear acrylamide as a carrier plus 10 μl of 5M NaCl and 1.1 ml of ethanol, and incubating 20 min at 0 °C. The DNA pellet was recovered by centrifugation, washed in 70% ethanol, dried under vacuum and resuspended in 50 µl of distilled water. The DNA (50 ng) was radioactively labelled and used to probe a cDNA library in λgt11.

The nitrocellulose filters were processed as described in Sambrook et al (1989). Duplicate filters were used, and the films were exposed 5-7 days in order to obtain a good signal. From among 200,000 plaques screened in this way, 31 hybridized to EW7D11. Among these 31 5 clones, 17 were homologous to each other, as checked by cross hybridization in stringent conditions. The size of the inserts in the 17 clones was estimated and the clone with the largest cDNA was retained for further analysis. A small scale preparation of this phage was prepared using the lambdasorb method, and the insert was excised by restricting 10 with EcoR1. This insert was ligated into a pBLUESCRIPT II vector linearized with EcoRI, and the ligation mixture was used to transform E. coli strain DH5α.

One of the recombinant clones was designated pBNDES3 (Figure 2), and retained for sequencing. The sequence was determined on 15 both strands, using the sequenase enzyme, (US Biochemicals, Cleveland, OH) according to the instructions provided by the supplier. The nucleotide sequence of the insert in pBNDES3 is presented as Figure 3. The deduced amino acid sequence of the largest open reading frame in the nucleotide sequence is also shown in Figure 3.

Comparison of the deduced amino acid sequence of the 383 amino acid open reading frame in clone pBNDES3 against the known sequences in GenBank release 70 was performed using the FASTA program (Lipman and Pearson, 1985). This analysis revealed that the sequence from pBNDES3 had a region of significant homology to a 25 previously characterized desaturase gene from the cyanobacterium Synechocystis (Figure 4). (Wada et al. 1990). This was considered suggestive evidence that the clone pBNDES3 encoded a desaturase which was probably the fad3 structural coding sequence product. This was subsequently confirmed by a genetic complementation experiment.

The cDNA was cloned into plant transformation vector pBI121 (Figure 5) under the control of the CaMV35S promoter to construct pTiDES3 (Figure 6). Plasmid pTiDES3 was introduced into an Agrobacterium tumefaciens strain which also carried an Ri plasmid and this 5 was used to produce transgenic rooty tumors from both wild type Arabidopsis and the fad3 mutant. Transgenic tissue was selected for antibiotic resistance to confirm the presence of the pTiDES3. Fatty acid methyl esters were then prepared and examined by gas chromatography to determine the profile of fatty acids being produced in the tissue. The levels of linolenic acid increased, demonstrating that the cDNA on pTiDES3 can complement the fad3 mutation. These results, which are described in detail in Example 1 below, confirm the identity of the cDNA as encoding a linoleic acid desaturase.

The isolation of a plant structural coding sequence provides those skilled in the art with a tool for the manipulation of gene expression by the mechanism of antisense RNA. The technique of antisense RNA is based upon introduction of a chimeric gene which will produce an RNA transcript that is complementary to a target gene (reviewed in Bird and Ray, 1991). The resulting phenotype is a reduction in the gene product from the endogenous gene. The portion of the gene which is sufficient for achieving the antisense effect is variable in that numerous fragments or combinations thereof are likely to be effective. Various portions of the structural coding sequence of linoleic acid desaturase isolated either from cDNA or genomic clones are likely capable of reducing linolenic acid levels in plants by reduction in levels of linoleic acid desaturase levels. An example of using an antisense oriented linoleic acid desaturase structural coding sequence is set out in Example 2.

Polyadenylation Signal

The 3' non-translated region of the double stranded DNA molecule of the present invention contains a region that functions in plant cells to promote polyadenylation to the 3' end of the RNA sequence. Any such regions can be used within the scope of the present invention. Examples of suitable 3' regions are (1) the 3' transcribed, non-translated regions containing the polyadenylated signal of Agrobacterium tumorinducing (Ti) plasmid genes, such as the nopaline synthase (NOS) gene, and (2) 3' regions of plant genes like the soybean storage protein genes and the small subunit of the ribulose-1,5-bisphosphate carboxylase (ssRUBISCO) gene. An example of a preferred 3' region is that from the NOS gene, described in greater detail in the examples below.

Plant Transformation/Regeneration

Any plant which can be transformed to contain the doublestranded DNA molecule of the present invention are included within the
scope of this invention. Preferred plants which can be made to have
increased or decreased linolenic acid content by practice of the present
invention include, but are not limited to sunflower, safflower, cotton, corn,
wheat, rice, peanut, canola/oilseed rape, barley, sorghum, soybean, flax,
tomato, almond, cashew and walnut.

A double-stranded DNA molecule of the present invention containing the functional plant linoleic acid desaturase gene can be inserted into the genome of a plant by any suitable method. Suitable plant transformation vectors include those derived from a Ti plasmid of Agrobacterium tumefaciens, as well as those disclosed, e.g., by Herrera-Estrella (1983), Bevan (1984), Klee (1985) and EPO publication 120,516 (Schilperoort et al.). In addition to plant transformation vectors derived from the Ti or root-inducing (Ri) plasmids of Agrobacterium, alternative methods can be used to insert the DNA constructs of this invention into

plant cells. Such methods can involve, for example, the use of liposomes, electroporation, chemicals that increase free DNA uptake, free DNA delivery via microprojectile bombardment, and transformation using bacteria, viruses or pollen.

A plasmid expression vector, suitable for the expression of the linoleic acid desaturase gene in monocots is composed of the following: a promoter that is specific or enhanced for expression in the lipid storage tissues and a 3' polyadenylation sequence such as the nopaline synthase 3' sequence (NOS 3'; Fraley et al., 1983). This expression cassette may be 10 assembled on high copy replicons suitable for the production of large quantities of DNA.

A particularly useful Agrobacterium-based plant transformation vector for use in transformation of dicotyledonous plants is plasmid vector pMON530 (Rogers, S.G., 1987). Plasmid pMON530 (see Figure 7) is a 15 derivative of pMON505 prepared by transferring the 2.3 kb StuI-HindIII fragment of pMON316 (Rogers, S.G., 1987) into pMON526. Plasmid pMON526 is a simple derivative of pMON505 in which the SmaI site is removed by digestion with XmaI, treatment with Klenow polymerase and ligation. Plasmid pMON530 retains all the properties of pMON505 and the 20 CaMV35S-NOS expression cassette and now contains a unique cleavage site for SmaI between the promoter and polyadenylation signal.

Vector pMON505 is a derivative of pMON200 (Rogers, S.G., 1987) in which the Ti plasmid homology region, LIH, has been replaced with a 3.8 kb HindIII to SmaI segment of the mini RK2 plasmid, pTJS75 25 (Schmidhauser & Helinski, 1985). This segment contains the RK2 origin of replication, oriV, and the origin of transfer, oriT, for conjugation into Agrobacterium using the tri-parental mating procedure (Horsch & Klee, 1986). Plasmid pMON505 retains all the important features of pMON200 including the synthetic multi-linker for insertion of desired DNA fragments,

the chimeric NOS/NPTII'/NOS gene for kanamycin resistance in plant cells, the spectinomycin/streptomycin resistance determinant for selection in E. coli and A. tumefaciens, an intact nopaline synthase gene for facile scoring of transformants and inheritance in progeny and a pBR322 origin of replication for ease in making large amounts of the vector in E. coli. Plasmid pMON505 contains a single T-DNA border derived from the right end of the pTiT37 nopaline-type T-DNA. Southern analyses have shown that plasmid pMON505 and any DNA that it carries are integrated into the plant genome, that is, the entire plasmid is the T-DNA that is inserted into the plant genome. One end of the integrated DNA is located between the right border sequence and the nopaline synthase gene and the other end is between the border sequence and the pBR322 sequences.

When adequate numbers of cells (or protoplasts) containing the linoleic acid desaturase gene are obtained, the cells (or protoplasts) are regenerated into whole plants. Choice of methodology for the regeneration step is not critical, with suitable protocols being available for hosts from Leguminosae (alfalfa, soybean, clover, etc.), Umbelliferae (carrot, celery, parsnip), Cruciferae (cabbage, radish, rapeseed, etc.), Cucurbitaceae (melons and cucumber), Gramineae (wheat, rice, corn, etc.), Solanaceae (potato, tobacco, tomato, peppers) and various floral crops. See, e.g., Ammirato (1984); Shimamoto, 1989; Fromm, 1990; Vasil and Vasil, 1990.

Uses of Linoleic Acid Desaturase

The present invention can be used for any modification (either increase, decrease, or mere change) of the oil content of a plant or plant tissue. Linolenic acid is an important constituent of several membranes in plant cells.

One preferred method is to modify the oil content of the plant to improve the plant's temperature sensitivity. For instance, plants deficient in linolenic acid display reduced fitness at low temperature (Hugly and Somerville, 1992). Also, increased linoleic acid content in vegetative tissues has been implicated as a factor in freezing tolerance in higher plants (Steponkus et al., 1990 and references therein). In a preferred embodiment, expression of the linoleic acid desaturase structural coding sequence can result in the genetic modification of higher plants to achieve tolerance to low environmental temperatures. Transformation with pTiDES3 demonstrates that linolenic acid levels can be increased by expression of this gene in a constitutive manner. Chilling or freezing injury in crops may be overcome by expression of this gene in vegetative or reproductive tissues by employing an appropriate promoter.

Linolenic acid, a polyunsaturated fatty acid, is also extensively used in the paint and varnish industry in view of its rapid oxidation. Flax seed is a predominant source of this oil. Higher quantities of this fatty acid in rapeseed or soybean will provide opportunities for using vegetable oils from these sources as a replacement for linseed (flax) oil. Expression of a linoleic acid desaturase structural coding sequence in seed tissue can result in a higher proportion of linolenic acid in the storage oil.

Linolenic acid is further a precursor in the biosynthesis of jasmonic acid, an important plant growth regulator. Linolenic acid is converted to jasmonic acid by introduction of an oxygen to the carbon chain by a lipoxygenase, followed by dehydration, reduction, and several β-oxidations (Vick and Zimmerman, 1984). The activity of jasmonic acid has been measured in terms of induction of pathogen defense responses. By application of free linolenic acid to plants, plant pathogen defenses can also be induced (Farmer and Ryan, 1992). A model has been proposed to explain the ability of free linolenic acid to exhibit the effects associated with jasmonic acid (Farmer and Ryan, 1992). It is hypothesized that all of the enzymatic activities which are required for the conversion of linolenic acid to jasmonic acid are constitutively present in the cell and the rate limiting

step in the production of jasmonic acid is the availability of free linolenic acid. A likely route for the production of the free linolenic acid is by the activity of a lipase in the plasma membrane.

It further has been observed that exogenous jasmonic acid can
more powerfully activate defense responses than can wounding. This
suggests that wounds cannot generate enough free linolenic acid to support
high level production of jasmonic acid. The activity of the lipase or the
availability of appropriate substrate for the lipase may be rate limiting
upon wounding. By increasing levels of available substrate, increasing
linolenic acid levels in the plasma membrane, it should be possible to
enhance a plant's ability to respond to pathogens by allowing for a higher
production of jasmonic acid. Expression of a linoleic acid desaturase
structural coding sequence can result in a higher molar percent linolenic
acid in the plasma membrane of a plant cell therefore enhancing the
jasmonic acid signaling pathway. It is our intent to evaluate plants
containing high linolenic acid levels in root and foliar tissues for their
pathogen resistance.

It is also undesirable to have significant levels of linolenic acid in cooking oils. Linolenic acid is unstable during cooking and is rapidly oxidized.

The oxidized products impart rancidity to the finished product. Rapeseed or soybean oil containing less than about 3%, and preferably 2% or less of linolenic acid is ideal for use as a cooking oil. By expression of the antisense of the structural coding sequence for linoleic acid desaturase, it is possible to reduce the linolenic acid content of these oils.

All higher plants have linolenic acid and, therefore, contain genes for linoleic acid desaturases. Because of the many examples in which genes isolated from one plant species have been used to isolate the homologous genes from other plant species, it is apparent to any one skilled in the art, that the results presented here do not only pertain to the use of the B.

napus fad3 gene, or to the use of the gene to modify fatty acid composition in B. napus. Obviously, the linoleic acid desaturases from many organisms could be used to increase linolenic acid biosynthesis and accumulation in plants and enzymes from any other higher plant or algae can serve as sources for linoleic acid desaturase genes. For example, since a YAC containing the Arabidopsis gene was used to isolate the B. napus gene, it is apparent that the insert in pBNDES3 could be used as a probe of genomic libraries for isolation of the corresponding full length genes from other plant species. It is also likely that the information contained in the sequence of this gene will be useful to clone other lipid desaturases genes.

Expression of a linoleic acid desaturase in a sense orientation may also allow for the isolation of plants with reduced levels of linolenic acid. This could be accomplished by the mechanism of co-suppression (Bird and Ray, 1991). The molecular mechanism of co-suppression is at this time poorly understood but occurs when plants are transformed with a gene that is identical or highly homologous to an allele found in the plants genome. There are several examples where expression of a chimeric gene in plants can result in a reduction of the gene product from both the chimeric gene and the endogenous gene(s). Those skilled in the art will recognize that the resulting decrease in linolenic acid would be a direct result of expression of the linoleic acid desaturase structural coding sequence and would be correlated to the linoleic acid desaturase activity in the transformed plant.

Linolenic acid levels in plant cells can also be modified by isolating genes encoding transcription factors which interact with the upstream regulatory elements of the plant linoleic acid desaturase gene(s). Enhanced expression of these transcription factors in plant cells can effect the expression of the linoleic acid desaturase gene. Under these conditions, the increased or decreased linolenic acid content would also be caused by a corresponding increase or decrease in the activity of the linoleic acid

desaturase enzyme although the mechanism is different. Methods for the isolation of transcription factors have been described (Katagiri, 1989).

The following examples are provided to better elucidate the practice of the present invention and should not be interpreted in any way to limit the scope of the present invention. Those skilled in the art will recognize that various modifications, truncations, etc. can be made to the methods and genes described herein while not departing from the spirit and scope of the present invention.

Example 1

10 Expression of fad3 gene to increase linolenic acid

To verify the assumption that the cDNA insert in pBNDES3 encodes a linoleic acid desaturase, both wild type and fad3 mutation Arabidopsis were transformed to contain the cDNA insert. In order to express the linoleic acid desaturase structural coding sequence (hereafter 15 referred to as the "fad3 gene") in plant cells, the plasmid pBNDES3 was digested with XhoI and the ends were filled in with the Klenow fragment of DNA polymerase (Sambrook et al 1989). The cDNA insert was subsequently excised by digestion with Sac1 and ligated into the Sac1 and Smal sites of the binary Ti plasmid vector pBI121 (Clontech 20 Laboratories), thereby replacing the GUS reading frame. The ligation reaction was carried out in 20 µl for 12 h at 16 °C using 100 ng of both insert and vector, and one unit of T4 DNA ligase. The ligation mixture was used to transform competent DH5\alpha E. coli cells (prepared by the calcium chloride method, according to Sambrook et al, 1989), and transformants 25 were selected on L-broth plates that contained 50 μg/μl Kanamycin. Alkaline minipreparations of recombinant clones were analyzed for the correct restriction pattern. One of these plasmids, designated pTiDES3, was used for further experiments.

This plasmid was electroporated (according to Mersereau and Pazour, 1990) into Agrobacterium tumefaciens strain R1000 which carries an Ri plasmid. The transformed bacteria were selected on kanamycin LB plates for 2 days at 30 °C. DNA minipreparations of several recombinant bacteria were performed and analyzed as described above to verify the presence of the construct.

Young flowering stems of wild type and the fad3 mutant of Arabidopsis were sterilized for 30 min in 10% commercial bleach, 0.02% Triton X100, and 2-cm explants that contained the flowering stem were infected with R1000 (pTiDES3) This was performed by dipping the sectioned extremity in a drop of an overnight culture of the appropriate Agrobacterium that was grown from a single colony in LB medium supplemented with 50 ug/ml Kanamycin.

The infected stems were cultured for two days on solid MSO medium (Gibco MS salts plus Gamborg B5 vitamins, 3% sucrose and 0.8% agar). At this time the stem segments were transferred for 5 weeks to MSO medium containing 200 µg/ml cefotaxime to kill the bacterium. After approximately two weeks, most of the stem explants had developed rooty tumors resulting from transfer of parts of the Ri plasmid into cells of the stem explants. In order to identify the rooty tumors which had also received the binary Ti plasmid pTiDES3, approximately 24 rooty tumors from each treatment were transferred to MSO medium containing 50 µg/ml of kanamycin to select for the growth of those roots which had been cotransformed with the binary Ti plasmid; the medium contained also 200 µg/ml of cefotaxime to inhibit bacterial growth. Following a further period of growth for 2 weeks, fatty acid methyl esters were prepared (as described above) from the roots for analysis by gas chromatography. The results of these analyses are presented in Table 2.

Table 2. Genotype

	mol% Fatty acid	wildtype pBI121	fad 3 pBI121	wildtype pTiDES3	fad3 pTiDES3
5					,
	16:0	22.0±2.9	21.2±1.6	21.1±0.9	21.3±2.3
	16:1	2.5±0.7	1.6±0.8	2.0 ± 0.1	1.5±0.2
	18:0	2.3±1.9	2.3±1.9	1.9 ± 0.2	1.6±0.4
	18:1	3.8±1.3	5.9 ± 2.6	7.7 ± 2.0	9.1 ± 2.0
10	18:2	37.3±3.7	62.2±5.9	15.7±11.7	24.4±14.9
	18:3	31.9±4.5	6.7±0.7	51.3±10.9	42.1±15.5

Table 2 shows the fatty acid composition of transgenic roots. The transgenic roots resulting from infection of wild type or the fad3 mutant with A. tumefaciens R1000 carrying the vector (pBI121) or the plasmid pTiDES3 were grown in the presence of kanamycin (50 g/ml) for three weeks to identify the roots which had been cotransformed with one of these plasmids. The fatty acid composition of the roots was determined as previously described (Browse et al., 1986). The abbreviations used in Table 2 are as follows: 16:0, palmitic acid; 16:1, palmitoleic acid; 18:0, stearic acid; 18:1, oleic acid; 18:2, linoleic acid; 18:3, linolenic acid. The values presented are the mean ± SD (n=12).

From these results it can be seen that the production of rooty tumors containing pBI121 on wild type Arabidopsis or the fad3 mutant had no effect on the fatty acid composition over non-pBI121 containing wild type Arabidopsis or fad3 mutant. By contrast, transformation of the fad3 mutant with the plasmid pTiDES3 resulted in large increases in the content of linolenic acid. In contrast to the linolenic acid content of 6.7 +/-0.7% in the fad3 mutant transformed with pBI121, the presence of pTiDES3 resulted in accumulation of 42.1% of the fatty acids as linolenic acid. The increased content of linolenic acid was accompanied by a

decrease of corresponding magnitude in the content of linoleic acid. Thus, it is clear that the fad3 gene encodes a linoleic acid desaturase. Introduction of the fad3 gene into wild type tissues also resulted in significantly increased accumulation of linolenic acid and a corresponding decrease in linoleic acid (Table 2). Thus, it is apparent from these results that the linoleic acid content of plant tissues can be increased by high level expression of a linoleic acid desaturase. In the present embodiment, the fad3 gene was placed under transcriptional control of the constitutive high level CaMV 35S promoter carried on pBI121. The implication from these results is that expression from this promoter raised the level of expression of the fad3 gene to levels higher than are normally achieved by expression from the endogenous fad3 promoter. The results presented here indicate that the fad3 gene has significant utility in genetic modification of higher plants to elevate linolenic acid levels.

15 Example 2

Antisense expression of fad3 gene to decrease linolenic acid levels

In order to decrease the linoleic acid desaturase activity by genetic engineering methodology, the cDNA insert of pBNDES3 was cloned into plant expression cassettes in an antisense orientation. A 959bp BgIII restriction fragment of pBNDES3 was used in the antisense expression vectors. The fragment is from 152 nucleotides downstream of the initiating methionine codon of the cDNA to a second BgIII restriction site that is located near the C-terminus of the coding region. 189 nucleotides of the coding region are excluded from this fragment. Triple ligations were performed with the fad3 gene fragment to construct two separate plant expression cassettes.

A seed specific expression cassette was constructed by insertion of the BgIII fragment of pBNDES3 in an antisense orientation behind the soybean promoter for the α ' subunit of β -conglycinin (7S promoter). A

975bp HindIII to BglII fragment containing the 7S promoter derived from pMON529 was prepared by digesting with BglII for 30min at 37 °C followed by addition of Calf Intestinal Alkaline Phosphatase (CIAP) (Boehringer Mannheim). The reaction was allowed to proceed for 20min followed by 5 purification of the linearized DNA using the GeneClean (Bio 101) purification system. The DNA was then digested with HindIII. A fragment derived from pMON999 containing the Nopaline synthase 3' region and the pUC vector backbone was prepared by digestion with BamHI and treatment with CIAP. The DNA was purified by the GeneClean procedure 10 and digested with HindIII. The fragment of pBNDES3 was prepared by digestion with BglII. The three fragments were purified by agarose gel electrophoresis and the GeneClean procedure. 50 to 200ng of the purified fragments were ligated for one hour at room temperature followed by transformation into the E. coli strain JM101. Resulting transformant 15 colonies were used for plasmid preparation and restriction digestion analysis. Double digestion with BglII and NcoI was used to screen for transformants containing the fad3 gene in an antisense orientation. One clone was designated as correct and named pMON13801.

A second expression cassette was constructed to allow for constitutive expression of the antisense message in plants. A fragment containing the enhanced 35S promoter was prepared from pMON999 by restriction digestion with HindIII and BglII followed by treatment with CIAP as above. The correct sized fragment was obtained by agarose gel electrophoresis and the GeneClean procedure. The BglII to HindIII vector fragment and the BglII fragment of pBNDES3 which were purified above were used in this construction. Ligation, transformation and screening of clones were as described above. One clone was designated as correct and named pMON13802.

In both pMON13801 and pMON13802, the promoter, fad3 gene and the Nos 3' region can be isolated on a Not! restriction fragment. These fragments can then be inserted into a unique NotI site of the vector pMON17227 to construct glyphosate selectable plant transformation 5 vectors. The vector DNA is prepared by digestion with NotI followed by treatment with CIAP. The fad3 containing fragments are prepared by digestion with NotI, agarose gel electrophoresis and purification with GeneClean. Ligations are performed with approximately 100ng of vector and 200ng of insert DNA for 1.5 hours at room temperature. Following 10 transformation into the E. coli strain LE392, transformants were screen by restriction digestion to identify clones containing the fad3 expression cassettes. Clones in which transcription from the fad3 cassette is in the same direction as transcription from the selectable marker were designated as correct and named pMON13804 (FMV/CP4/E9, 7S/anti fad3/NOS) 15 (Figure 8) and pMON13805 (FMV/CP4/E9, E35S/anti fad3/NOS) (Figure 9).

In preparation for transforming canola cells, pMON13804 and pMON13805 were mated into Agrobacterium ABI by a triparental mating with the helper plasmid pRK2013.

Seeds from the plants produced by transformation were analyzed for alterations in fatty acid profile. Fatty acid methyl esters (FAMES) were prepared from seed tissue and analyzed by capillary gas chromatography (Browse et al, 1986). For initial screening of plants, six seeds were pooled together from an individual plant. The seeds were 25 crushed and FAMES extracts were made. Control plants, plants transformed with the selectable marker only (pMON17227), were also analyzed using the identical procedure. From the initial screen on pooled seed samples, several lines were identified which displayed a decreased level of linolenic acid. Lines with decreased levels of linolenic acid were

reanalyzed by determining fatty acid profiles from individual seeds. Four to twenty individual seed were analyzed from candidate lines and from selected control plants. The results of the FAMES analysis is summarized in Figure 9.

Figure 9 shows the levels of fatty acids expressed in molar percent of twenty individual seed of the transgenic line 13804-51 as compared to control seed. Panel A discloses oleic acid, panel B discloses linoleic acid and panel C discloses linolenic acid.

The data in Figure 9 demonstrate that antisense expression of a linoleic acid desaturase has significantly altered the fatty acid profile of the resulting seed tissue. The percent of linolenic acid has been reduced to a little over 2% of the total fatty acid in the seed tissue. The percent of linoleic acid has been reduced slightly and surprisingly, the percent of oleic acid in the seed has been increased to approximately 70%. This demonstrates the applicability of utilizing the fad3 gene to manipulate the fatty acid profile of crop plants.

In order to demonstrate that the alteration in the fatty acid profile of the FAMES extracted from total seed tissue would be reflected in the seed oil fraction, triglycerides from seeds of fad3 antisense plants were characterized. Total lipid extracts were made by pooling ten seeds and grinding in 2ml of methanol:chloroform:water (4:2:1). The homogenate was allowed to stand for 20min and then debris was pelleted and discarded. To the supernatant 400µl of chloroform:methanol (2:1), 640µl of chloroform and 740µl of water was added and vortexed. Phases were separated by centrifugation and the chloroform phase was recovered and dried under nitrogen. Samples were resuspended in 100µl of chloroform and 10µl was applied to silica gel G thin layer chromatography plates for separation. Two identical plates were prepared with one being charred after development to allow for alignment and location of spots to be analyzed on

the other plate. Plates were developed three times in petroleum ether:diethyl ether:acetic acid (90:10:1). One plate was sprayed with 50% sulfuric acid and heated in an oven at 90°C to allow for detection of lipids. Triglyceride fractions were identified as comigrating on the plate with purchased lipid standards (Sigma Chemical Co, cat #178-13). The charred plate was aligned with the identical plate and the triglyceride fractions were scraped from the plate. The fatty acids were transesterified to produce FAMES extracts for GC analysis by the same procedure as above. The fatty acid profiles of the triglyceride fractions are shown in Table 3 and demonstrate that this fraction have decreased linolenic acid.

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15	Transgenic <u>line</u>	Mol% 18:1	<u>18:2</u>	18:3
-	17227-10	44	30	15.3
	17227-493	65	17	6.9
	13804-47	58	21	4.3
20	13804-50	67	20	2.8
	13804-76	59	19	5.0
	13804-117	62	21	4.0

Table 3 compares the fatty acid molar percentages of triglyceride fractions from control and transgenic lines. These above results provide clear evidence that the fad3 gene can be used to decrease the levels of linolenic acid in the storage oil of plants. The gene provides a tool for the manipulation of the fatty acid profile of seed storage oil to improve the products derived from the oil.

A surprising result of this Example 2 is the effect the antisense fad3 gene has on the oleic acid content. The precise mechanism by which

antisense expression of a gene exerts an effect on the activity of an endogenous gene is unclear but is obviously a function of the homology of the sense and antisense gene products. Based upon the above experimental result, it would not be unreasonable to predict that the 5 portion of the fad3 gene antisense message used contained a certain degree of homology with the genes providing the activity of one or more oleate desaturases. Therefore, a further advantage of the above invention is that it is possible that expression of a linoleic acid desaturase antisense message may exert an effect on oleate desaturase activity.

The unexpected nature of the reduction in oleic acid desaturase activity from the antisense fad3 plants is most apparent when one compares the fatty acid profiles from the antisense plants and the fad3 mutant of Arabidopsis. The levels of linoleic acid in the fad3 mutant plants increased when linoleic acid desaturase activity was eliminated by 15 mutation. This indicates that the activity of the oleate desaturase was not highly effected by the loss of linoleic acid desaturase activity or by the accumulation of linoleic acid. In the fad3 mutant of Arabidopsis the level of linoleic acid increased when the level of linolenic acid decreased. However, a different pattern occurred in the antisense fad3 plants. In plants which 20 exhibit a decreased percent of linolenic acid there is no corresponding increase, and is often a decrease, in the percent of linoleic acid. There is an increase in the percent of oleate in the antisense fad3 plants. This would indicate that oleate desaturase activity is depressed in these plants. The effects on the fatty acid profile by the fad3 mutation and the fad3 antisense 25 expression are not equivalent, indicating that antisense expression of a linoleic acid desaturase can depress an oleate desaturase activity in plants.

Example 3

Modification of linolenic acid levels in soybean

The isolation of the fad3 gene from B. napus provides a tool to those with ordinary skill in the art to isolate the corresponding gene or cDNA from other plant species. There are many examples in which genes from one plant species have been used to isolate the homologous genes from another plant species. One such plant which could be improved upon by the modification of the level of linolenic acid is soybean.

Soybean oil typically contains linolenic acid at a level of 7-9% of the fatty acid in the oil. This level is undesirable because it promotes instability upon heating and imparts rancidity to the finished product. The levels of linolenic acid can be lowered by the expression of the soybean fad3 gene or cDNA in an antisense orientation in the developing seed. The following example describes one method for the isolation of a fad3 cDNA from soybean. However, similar procedures could be followed to isolate a genomic clone which could also be used to decrease the level of linoleic acid desaturase activity by antisense expression of a portion or all of the gene.

The fad3 gene from B.napus is used as a probe to screen a cDNA library constructed from soybean mRNA. In order to isolate a cDNA to be used in decreasing linolenic acid in seed, the optimal tissue to use for the isolation of mRNA is developing seed. There is, however, flexibility in the choice of methods and vectors which can be used in the construction and analysis of cDNA libraries (Sambrook et al, 1989). Procedures for the construction of cDNA libraries are available from manufacturers of cloning materials or from laboratory handbooks such as Sambrook et.al, 1989. Once a suitable cDNA library has been constructed from soybean, all or a portion of the fad3 cDNA from B.napus is labeled and used as a probe of the library. DNA fragments can be labeled for radioactive or non-radioactive screening procedures. The library is screened under suitable stringency.

Conditions are dependent upon the degree of homology between the fad3 gene of B. napus and soybean. Probe positive clones are plaque purified by standard procedures and characterized by restriction enzyme mapping and DNA sequence analysis. Clones are concluded to be soybean fad3 based upon data obtained from the sequence analysis or by expression in plants.

The entire clone or a portion thereof is placed down stream of a promoter sequence in an antisense orientation. Suitable promoters include seed specific promoters, such as the 7S (β-conglycinin) α'-subunit promoter, or less tissue specific promoters, such as the CaMV 35S 10 promoter. An appropriate 3' non-translated region is placed downstream of the antisense cDNA to allow for transcription termination and for the addition of polyadenylated nucleotides to the 3'end of the RNA sequence. This expression cassette is then combined with a selectable or scorable marker gene and soybean cells are transformed by free DNA delivery 15 (Christou et al, 1990) or an Agrobacterium based method of plant transformation (Hinchee et al, 1988). Plants recovered are allowed to set seed and mature seed are used for the production of FAMES by the procedures outlined above. The FAMES extracts are analyzed by gas chromatography to identify plant lines with reduced levels of linolenic acid in the seed. 20

Alternatives to the above methods may include but are not limited to the use of degenerate oligonucleotides as probes to screen the library. Degenerate oligonucleotide probes would be most optimally designed by choosing short segments of the fad3 amino acid sequence where the degeneracy of the genetic code is limited or by choosing sequences which appear to be highly conserved between the fad3 gene of B. napus and other known linoleic acid desaturases, such as the desaturase from the cyanobacterium Synechocystis. The oligonucleotides could be labeled and used to probe a soybean cDNA library. Alternatively, degenerate

oligonucleotides could be used as primers for the isolation of a portion or all of the soybean cDNA by PCR amplification.

Similar procedures could be used to isolate the homologous genes from other plant species. Another preferred plant species which could be improved upon by the modification of the level of linolenic acid is flax. Flax oil typically contains linolenic acid at a level of 45-65% of the fatty acid in the oil. This level is undesirable because it promotes instability upon heating and imparts rancidity to the finished product.

Example 4

10 Sense expression of fad3 to obtain reduced levels of linolenic acid

The cloning of the fad3 gene also provides a tool to decrease the levels of linolenic acid via the mechanism of co-suppression. The molecular mechanism of co-suppression occurs when plants are transformed with a gene that is identical or highly homologous to an allele found in the plants genome (Bird and Ray, 1991). There are several examples where expression of a chimeric gene in plants can result in a reduction of the gene product from both the chimeric gene and the endogenous gene(s). Therefore the fad3 gene product of B. napus may be reduced by transformation of B. napus with all or a portion of the fad3 cDNA which has been isolated. The resulting plant has reduced linoleic acid desaturase activity in tissues where the chimeric gene is expressed. The phenotype of reducing the linoleic acid desaturase activity is a reduction in the levels of linolenic acid. The mechanism of co-suppression could be applied to any plant species from which the fad3 gene is cloned and the plant species is transformed with fad3 in a sense orientation.

In order to reduce levels of linolenic acid by the mechanism of cosuppression, a plant transformation construct is assembled with the fad3 gene or cDNA in a sense orientation. The entire clone or a portion thereof is placed downstream of a promoter sequence in a sense orientation. Suitable promoters include seed specific promoters, such as the 7S (β-conglycinin) α'-subunit promoter, or less tissue specific promoters, such as the CaMV 35S promoter. An appropriate 3' non-translated region is placed downstream of the fad3 gene to allow for transcription termination and for the addition of polyadenylated nucleotides to the 3' end of the RNA sequence. This expression cassette is then combined with a selectable marker gene and B. napus cells are transformed by an Agrobacterium based method of plant transformation. Plants recovered are allowed to set seed and mature seed are used for the production of FAMES which are analyzed by gas chromatography to identify plant lines with reduced levels of linolenic acid in the seed.

Example 5

Isolation of a chloroplast delta 15 desaturase from Arabidopsis

A fragment of 959bp was excised from the fad3 cDNA insert 15 using the restriction endonuclease BglII, and labeled radioactively according to Feinberg and Vogelstein (1983). This fragment was used to probe a cDNA library from Arabidopsis thaliana as described above (Example 1) except that the hybridization temperature was 52° C. Several cDNA clones were positive, and one of them (pVA1) was further characterized. 20 Its deduced amino acid sequence exhibited a strong homology with fad3 except at the N-terminus. The cDNA insert was placed under the control of the 35S promoter in the Ti vector pBI121, and the resulting construct, pBIVA12 was electroporated into Agrobacterium (C58 pGV3101). The bacterium was used to transform the Arabidopsis mutant fadD. For 25 transformation, plants were grown at 22° C with a light intensity of 100/μE/cm-2, until bolting (approximately 2 and 1/2 weeks). The stems (1mm-10mm long) were removed and the plants were inoculated with a drop of an overnight culture of the bacterium. The same operation was repeated 7 days afterwards.

The plants were then allowed to set seeds. The seeds were plated (2500 seeds per 150mm petri dish) on MSO plates that contained 50µg/ml kanamycin to select for plants that had integrated the construct. One transformant plant was obtained, and the fatty acids from its leaves were analyzed by gas chromatography (Table 4). The results obtained show that the pBIVA12 construct is able to reestablish the levels of linolenic and hexadecatrienoic acids in the fadD mutant at a level equal to or superior to the wild type. This demonstrates that pVA12 encodes the fadD gene.

10

TABLE 4

·	fatty acid	fadD	WT	FadD pBIVA12
15				
	16:0	13.0	14.0	14.9
	16:1	4.9	4.3	4.2
	16:2	8.7	0.5	0.3
	16:3	3.0	13.2	9.5
20	18:1	3.3	2.3	1.2
	18:2	36.4	10.9	5.8
	18:3	30.8	54.6	63.7

Table 4 shows the complementation of the fadD mutant.

25 Fatty acids were extracted from leaves of *Arabidopsis* according to Browse et al (1986) and were quantified (mol%) by gas chromatography. WT stands for the Columbia wild type.

Example 6

Isolation of a second chloroplast delta 15 desaturase from Arabidopsis

A fragment of 959 bp was excised from the cDNA insert using the restriction endonuclease BgIII, and labelled radioactively according to 5 Feinberg and Vogelstein (1983). This fragment was used to probe a cDNA library from Arabidopsis, exactly as described above (Example 5). Among the several positive clones obtained, the cDNA pVA34 was further characterized. Its deduced amino acid sequence exhibited 71.8% and 79.5% homology with fad3 and fadD, respectively. The N-terminus resembled a 10 chloroplast transit peptide, meaning that this protein is likely to be localized to the chloroplast. The strong homology with fad3 and fadD suggests that the protein is also a delta 15 desaturase. Aside from fad3 and fadD, the only locus known to control delta 15 desaturation is the fadE locus, which controls a temperature-induced delta 15 desaturase.

15 Therefore, it is likely that the cDNA contained within the clone pVA34 corresponds to the fadE locus.

Example 7

Linoleic desaturase homology to plant oleic desaturases

The linoleic desaturase genes are the first plant desaturases isolated whose proteins enzymatically perform the desaturation of an unsaturated fatty acid precursor. The reaction that linoleic desaturase performs and the cofactors it uses are likely to be very similar for the oleic desaturase reaction. Given the similar reactions, similar substrates and probably similar cofactors, it is likely that the oleic desaturase genes and proteins have homology to the linoleic desaturase genes and proteins. That the genes share homology is supported by the finding that antisense expression of the linoleic acid desaturase message results in higher oleic acids levels, which experimentally indicates homology between the linoleic and oleic desaturases. These factors indicate that the linoleic desaturase

protein and nucleic acid sequences provide useful information for isolating other lipid desaturase genes, particularly oleic desaturase genes.

a. <u>Identification of unknown cDNA sequences in databases.</u>

Random cDNA sequencing generates a large number of sequenced clones but provides no information about the function of the encoded proteins. Homology to known proteins is the quickest method for identifying the protein function encoded in the sequenced cDNA. However, homology searches are informative only when a homology with a previously characterized protein are found. A cDNA sequence that is not homologous to any known protein remains in the unknown function category. Thus the results functionally identifying the linoleic desaturases by sequence and by their ability to complement mutations in plant desaturase genes now provides a method for identifying the function and identity of random cDNA clones by their homology to the linoleic desaturases. Additionally oleic desaturases are identified by their homology with linoleic desaturases.

A TFASTA search of the GenBank and EMBL public data bases for genes encoding proteins homologous to the protein sequence of the linoleic desaturase fad3 has identified both linoleic desaturases and a second class of plant lipid desaturases likely to be oleic desaturases. In particular, sequences found in GenBank and EMBL and identified as T04093 and T12950 show significant homology to linoleic desaturases but show less homology than other linoleic desaturases. These sequences have 30% homology to fad3 and 56% similarity to fad3 linoleic desaturase (TABLE 5). The full length clone of these cDNAs is obtained by standard methods and is inserted into plant gene expression and transformation vectors and transformed into fad2 Arabidopsis mutants to confirm the identity of the oleic desaturase by genetic complemention as was described in the example with linoleic desaturase.

-48-

TABLE 5

	Compa	parison of Fad3 and T				T0409	3 Protei	n Sequenc	es
5									
	Percent	Similar	ity:	52.38	18	Percent	Identity:	30.476%	
	fad3 101	GHGSFS	DIPL	LNSVVG	HILHS	FILVPYHO	GWRISHRTHH	QNHGHVENDESI	√ 150
10	T04093				1:11	1:1111 :	:1:.111 11	: . SNTGSLERDEVI	
	151	VPLPEK	LYKNI	LP	HSTRI	MLRYTVPI	PMLAYPIYLI	WYRSPGKEGSHI	F 195
15	35	VPKQKS	AIKW	 YGKYLNI	NPLGR	:: .: IMMLTVQF	: :: : `.VLGWPLYL	:. :: :: :: :: :: :: :: :: :: :: :: :: :	: 7 80
	196	NPYSSL	FAPSI	ERKLIA:	Isticv	VSIMLATL	VYLSFLVDP	VTVLKVYGVPYI	245
20		: . : . :	1 1.		:			• • • • • • • • • • •	
							•		
	· ·								

b. Isolation of a oleic desaturase cDNA.

The protein sequence of plant linoleic desaturases can be used to isolate oleic desaturases. The conserved regions between the linoleic desaturases and the DesA oleic desaturase are functionally important and are conserved in the plant oleic desaturase proteins as well. These conserved amino acid sequences provide a method of isolating plant oleic 30 desaturases. There are several regions of the linoleic desaturase fad3 that are conserved in fadD, fadE and DesA. The consensus amino acid sequence is shown in Table 6, with the amino acids identical in all four proteins shown in capital letters. As described below, oligonucleotides designed to encode the amino acids sequences in the conserved regions are used to identify and 35 isolate plant oleic desaturases.

-49-

TABLE 6

Fad3 Protein Sequence and Peptide Targets

MVVAMDQRSNVNGDSGARKEEGFDPSAQPPFKIGDIRAAIPKHCWVKSPLRSMSYVTRD v.tplttp ...spseed..erfdpgapppf.laDIraaiPKhCwvKnpwksmsyVvrd DIraaiPKhCwvK (la) DIraaiP (1b) aiPKhC (1c) KhCwvK 10 IFAVAALAMAAVYFDSWFLWPLYWVAQGTLFWAIFVLGHDCGHGSFSDIPLLNSVVGHIL va.vfalaa.aayfnnW.lwPlyW.aqGTmfwalFVlGHDCGHgSFsndp.lNsvvGH.l FV1GHDCGHqSF WflwPlyWvagGT (2a) WflwPlyW (3a) FV1GHD (3b) VlGHDC 15 (2b) WflwP (3c) GHDCGH (2c) wPlyW (2d) WvagGT (3d) CGHgSF HSFILVPYHGWRISHRTHHONHGHVENDESWVPLPEKLYKNLPHSTRMLRYTVPLPMLAY $\verb|hssilvPyHgWRisHrtHHqnhghvEnDesWhPl.ekiyknlpk.trmfrftlpipmlay| \\$ 20 **PvHqWRisHrtHH EnDesWvP** (5a) EnDesW (4a) PyHgW (5b) DesWvP (4b) HaWRisH (4c) WRisHrtHH 25 (4d) WRisH (4e) HrtHH PIYLWYRSPGKEGSHFNPYSSLFAPSERKLIATSTTCWSIMLAT.LVYLSFLVDPVTVLK pfylw.rspgk.gShyhpds.lF.pkerkdvltStacwtamaAl.lvcLnft.gpiqmlK 30 VYGVPYIIFVMWLDAVTYLHHHGHDEKLPWYRGKEWSYLRGGL.TTIDRDYG.IFNNIH lygiPywifvmWldfvTylHHnghedklpwyrgkeWSylrggL.tTldrDYg.winnin WSylragL.tTidrDY WldavTylHH (6a) WldavT (7a) WSylrggL 35 (7b) L tTidrD (6b) TylHH (7c) TidrDY

HDIGTHVIHHLFPQIPHYHLVDATRAAKHVLGRYYREPKTSGAIPIHLVESLVASIK HDIgtHviHHLfpqIPhYhLveAteaaKpvlGkyyrEpk.sgplplhLlesl.ksik HDIgtHviHHLfpqIPhY

5 (8a) HDIgtH

(8b) HviHHL

(8c) HHLfpqI

(8d) HLfpqIP

(8e) LfpqIPhY

10

15

KDHYVSDTGDIVFYETDPDLYVYASDKSKIN*
.dhyvsdtGdvvyYeadp.lyg..s*

c. <u>Isolation of the fadC (fad6) Gene from Arabidopsis thaliana</u>

The fadC gene (also referred to as fad6) encodes a

chloroplastic omega-6 desaturase.

The deduced amino acid sequences of the fad3 gene from Brassica napus and the fadD and fadE genes from Arabidopsis thaliana were compared with the DesA gene from Synechocystis (Nature, 347:200, 1990). The sequence GHDCGH was determined to represent the most highly conserved region of these proteins. Consequently, a degenerate oligomer was designed that contains all the possible condons for the sequence GHDCGH. This oligomer has the following sequence: GGNCAYGAYTGYGGNCA.

An Arabidopsis thaliana cDNA phage library obtained from the laboratory of Dr. Ron Davis (*PNAS*, 88: 1731-1735) was used to screen for desaturase genes. This library was made using material from all above ground plant parts.

Approximately 120,000 phage from the library were plated onto three plates and hybondN+ was then used to prepare three filters from each plate (*Molecular Cloning - A Laboratory Manual*, 2nd Edition. Eds. J. Sambrook, E. F. Fritsch, and T. Maniatis, Cold Spring Harbor Laboratory

Press, Cold Spring Harbor, New York 1989, hereafter "Sambrook"). Two filters from each plate were probed using the degenerate consensus oligomer which had been end-labelled with (32)P using T4 polynucleotide kinase. The hybridizations were performed in a solution that contained high 5 amounts of tetramethylammonium chloride in order to minimize differences in the melting temperatures of the oligomers that together comprise the degenerate consensus oligomer. The hybridization solution had the following composition: 3 M tetramethylammonium chloride, 10 mM sodium phosphate pH 6.8, 1.25 mM EDTA, 0.5% SDS, 0.5% milk. Hybridization 10 was carried out overnight at a temperature of 44°C. Filters were then washed four times, 20 minutes each time, with 6 x SSC + 0.15% SDS at room temperature. Filters were then washed one time, for 30 minutes, with 4 x SSC + 0.1% SDS at room temperature. The filters were then exposed to film for two days.

The third set of filters that were made from each phagecontaining plate were probed using DNA sequences from the three Arabidopsis desaturase genes that had already been identified: fad3, fadD and fadE. The fad3, fadD and fadE genes were labelled with (32)P and hybridized to the third set of phage filters in the following hybridization 20 solution: 0.2 M NaCl, 20mM sodium phosphate pH 7.7, 2mM EDTA, 1% SDS, 0.5% milk, 10% dextran sulfate, 0.1% sodium pyrophosphate. Hybridization was carried out overnight at 65°C. Filters were washed four times, 30 minutes per time, in 2 x SSC + 0.15% SD at room temperature and then for 45 minutes with 1 x SSC + 0.1% SDS at 65° C. The filters were then exposed to film for approximately two hours.

The two sets of filters that were probed with the degenerate consensus oligomer showed about 60 positive phage per plate (or about 180 total positive phage). Results from the third set of filters that were probed with the fad3, fadD and fadE genes indicated that only a small percentage

of the phage that hybridized to the consensus of oligomer contained the fad3, fadD or fadE genes.

Seventy-six of the phage that hybridized to the consensus oligomer, but not to the fad3, fadD or fadE genes, were plaque purified. The 5 purified phage were then spotted onto bacteria growing on solid media on plates and allowed to form plaques. Several duplicate filters were then made of these plates (Sambrook). One of these filters was probed with the consensus oligomer, as described above. A second filter was probed with a mixture of the Arabidopsis thaliana fad3, fadD and fadE genes, as described above.

In order to determine which of the 76 phage contained the same cDNA inserts as which other phage, some of the filters were probed with cDNA inserts from some of the phage. In order to perform this experiment, the cDNA inserts from most of the phage were isolated by 15 using oligomers that bound to DNA flanking the cDNA cloning site in the phage vector to isolate the cDNA sequences using the polymerase chain reaction (PCR). These cDNA sequences were labelled with (32)P (random hexamer labelling) and hybridized to the filters using the following hybridization solution: 30% formamide, 0.2M NaCl, 20mM sodium 20 phosphate pH 7.7, 2mM EDTA, 1% SDS, 0.5% milk, 0.1% sodium pyrophosphate. The hybridizations were carried out for 14 hours at 65°C. The filters were washed four times 15 minutes per wash, with 2 x SSC + 0.15% SDS at room temperature and were then exposed to film.

The combination of the high formamide concentration in the 25 hybridization solution and the high hybridization temperature meant that only DNA sequences that were virtually identical would hybridize, allowing us to distinguish between nearly identical sequences. Several rounds of hybridizations using cDNA inserts from different phage were carried out until it had been determined which phage contained the same, or at least extremely similar, cDNA inserts. On the basis of these experiments, we determined that all of the 76 phage contained one of four cDNA inserts. Sequence data was obtained from each of these four cDNAs. None of these cDNAs was found to be homologous to known desaturase genes, and so we feel that none of these four cDNAs is likely to encode a desaturase.

Since the number of phage that hybridized to the consensus oligomer was quite high (about 180 phage hybridized in the initial screen described above), we were not able to analyze all of the positive phage in the initial experiments. So, an attempt was made to identify phage that 10 hybridized to the consensus oligomer but that did not contain the fad3, fadD of fadE genes or one of the four non-desaturase encoding clones that were identified in the first screen. In order to do this, between 500,000 and 1,000,000 phage from the library described above were plated onto 10 plates. Three filters were made from each plate (Sambrook). Two of these 15 three sets of filters were then hybridized with (32) P labelled consensus oligomer as described above except that hybridization was carried out at 42°C instead of at 44°C. The third set of filters were hybridized with (32)P labelled DNA from the Arabidopsis fad3, fadD and fadE genes together with DNA from each of the four cDNA's identified in the first round of screening 20 as hybridizing to the consensus oligomer but not encoding desaturases. This third set of filters were hybridized in: 30% formamide, 0.2 M NaCl, 20mM sodium phosphate pH 7.7, 2mM EDTA, 1% SDA, 0.5% milk, 0.1% sodium pyrophosphate at 65°C. All three sets of filters were hybridized for 12 hours and then washed several times with 2 x SSC + 0.15% SDS at 25 room temperature. The filters were then exposed to film.

Approximately 200 phage from each plate hybridized to the consensus oligomer. 50-60% of these phage also hybridized to fad3, fadD, fadE or to one of the four clones identified in the first screen. About 58 phage that hybridized to the consensus oligomer, but not to fad3, fadD,

fadE or one of the four previously identified clones, were plaque purified. The purified phage were then spotted onto a bacterial lawn growing on solid media on a petri plate and the phage were allowed to form plaques. Several filters were prepared from these plates and hybridized with (32)P labelled cDNA inserts from various of the newly purified phage, as described above. In this manner, all of the phage identified in this second round of screening were found to contain one of eight different cDNA inserts.

Sequence data was obtained from each of the eight cDNA's. One of the cDNA's, which was contained within only one of the phage, was found to have some sequence similarity of a known desaturase gene from cyanobacteria, the DesA gene. Further sequence information was obtained for this clone. This additional sequence showed very significant sequence similarity to the DesA gene, confirming that the clone contained a desaturase gene. The remainder of the cDNA contained within the clone was sequenced and compared with the sequences of other known desaturases. The new desaturase was 53.0% identical to DesA at the nucleotide level and 43.9%, 45.6% and 47.0% identical to B. napus fad3, Arabidopsis fadD and Arabidopsis fadE, respectively. As the gene contained within the clone was significantly more similar in sequence to the DesA gene (which is a delta-12 desaturase) than to fad3, fadD or fadE (which are omega-3 desaturases), the new desaturase was expected to be a delta-12 (= omega-6) desaturase.

The additional sequence data also indicated that this new desaturase gene contains a region that has only a one base pair mismatch to the desaturase consensus sequence described above. This mismatch means that the new desaturase has the sequence GHDCAH instead of GHDCGH.

A clone containing a full length cDNA for this gene was isolated and completely sequenced. This full length cDNA was sub-cloned

into the plant transformation vector pBII121 such that the gene is transcribed under the control of the 35S promoter. This construct was then used to complement the phenotype of a fadC mutant (*Plant Phys.* 90: 522-529, 1989) of *Arabidopsis thaliana*, indicating that the gene encodes a chloroplastic omega-6 desaturase.

d. Proposed isolation of fad2

The most highly conserved peptide regions in the linoleic desaturases and the DesA desaturase were chosen as regions likely to be conserved in oleic desaturases. These 8 conserved regions are shown in TABLE 6. These regions were chosen on the following basis: These regions have areas highly conserved between the 3 linoleic desaturases and DesA, with at least 4 identical amino acids over a 10 amino acid span. Once a region was identified as conserved, the fad3 linoleic desaturase sequence was used as the amino acid sequence for the source of homology to identify oleic desaturases. This is because both fad3 and the non-plastid oleic desaturases are thought to be localized to the endoplasmic reticulum and are most likely to contain similar amino acid sequences.

Several peptide endpoints in each conserved area were chosen as the basis to subsequently design oligonucleotide probes for identifying the oleic desaturase gene. The peptide endpoints were chosen to be between 5 and 9 amino acids in length. The peptide end points were chosen to end on the conserved (identical) amino acids, and most often to begin on conserved amino acids. The rationale is that within the larger conserved area, some amino acid portions are more highly conserved than others, that 15 to 27 (5 to 9 amino acids) nucleotides is a good primer size for PCR, and that for PCR it is important that the 3' end of the primer matches the target, with the conserved (identical) amino acids the most likely to be present in the oleic desaturases. These 28 "oleic desaturase" peptide targets (Table 6) are the basis oligonucleotides that are designed for

hybridizing to the oleic desaturase cDNA sequences to identify and isolate the oleic desaturase cDNA clone.

Several possible methods for designing oligonucleotides and isolating the genes encoding the target peptide regions are known. For a 5 discussion of designing degenerate oligonucleotides see PCR Protocols - A Guide to Methods and Applications, Eds M. A. Innis, D. H. Gelfand, J J Sninsky and T. J. White, Academic Press, San Diego, California, 1990; and The two most common screening methods using the oligonucleotides are screening cDNA libraries and PCR amplification of 10 specific cDNAs. Gene probes from fad3, fadD and fadE are used under stringent hybridization conditions to identify these cDNAs and discard them in the screen for oleic desaturase cDNA clones. The method for using degenerate oligonucleotides to screen a cDNA library has been described in the example above demonstrating the isolation of the fadC oleic desaturase 15 gene. An immature plant seed active in oil biosynthesis, generally 2 to 5 weeks after pollination, preferably about 3 to 4 weeks after pollination, of a plant such as Arabidopsis or canola is used as the source of mRNA for making cDNA. First strand cDNA is made from the isolated mRNA and hybridized under stringent conditions in solution to an excess of biotinylated 20 fad3, fadD and fadE cloned cDNAs. The hybrids and biotinylated nucleic acids are removed with strepavidin and a second round of substraction is done to remove any remaining fad3, fadD and fadE sequences. The cDNA remaining in solution is used for PCR reactions. (For 5' RACE, see below, a polyA tail is added to the first strand cDNA 3' end).

A method that can readily evaluate a number of degenerate oligonucleotides probes is degenerate PCR (See chapters by Compton and by Lee and Caskey in *PCR Protocols*, cited above). In this method a degenerate set of oligonucleotides encompassing all the possible codon choices for the target peptide is synthesized (such degenerate

targets (Table 6) are the basis oligonucleotides that are designed for hybridizing to the oleic desaturase cDNA sequences to identify and isolate the oleic desaturase cDNA clone.

Several possible methods for designing oligonucleotides and 5 isolating the genes encoding the target peptide regions are known. For a discussion of designing degenerate oligonucleotides see PCR Protocols - A Guide to Methods and Applications, Eds M. A. Innis, D. H. Gelfand, J J Sninsky and T. J. White, Academic Press, San Diego, California, 1990; and The two most common screening methods using the Sambrook. 10 oligonucleotides are screening cDNA libraries and PCR amplification of specific cDNAs. Gene probes from fad3, fadD and fadE are used under stringent hybridization conditions to identify these cDNAs and discard them in the screen for oleic desaturase cDNA clones. The method for using degenerate oligonucleotides to screen a cDNA library has been described in 15 the example above demonstrating the isolation of the fadC oleic desaturase gene. An immature plant seed active in oil biosynthesis, generally 1 to 5 weeks after pollination, preferably about 2 to 4 weeks after pollination, of a plant such as Arabidopsis or canola is used as the source of mRNA for making cDNA. First strand cDNA is made from the isolated mRNA and 20 hybridized under stringent conditions in solution to an excess of biotinylated fad3, fadD and fadE cloned cDNAs. The hybrids and biotinylated nucleic acids are removed with strepavidin and a second round of substraction is done to remove any remaining fad3, fadD and fadE sequences. The cDNA remaining in solution is used for PCR reactions. (For 5' RACE, see below, a 25 polyA tail is added to the first strand cDNA 3' end).

A method that can readily evaluate a number of degenerate oligonucleotides probes is degenerate PCR (See chapters by Compton and by Lee and Caskey in *PCR Protocols*, cited above). In this method a degenerate set of oligonucleotides encompassing all the possible codon

-58-

TABLE 7
Peptide Targets for Fad2 Cloning

5	Peptide	sequence	Oligo sequence 5' - 3'
	1a	DIRAAIP	GAYATHMGNGCNGCNATHCC
	1b	AIPKHC	GCNATHCCNAARCAYTG
	1c	KHCWVK	AARCAYTGYTGGGTNAA
,	2a	WFLWPLYW	TGGTTYYTNTGGCCNYTNTAYTGG
10	2b	WFLWP	TGGTTYYTNTGGCCN
	2c	WPLYW	TGGCCNYTNTAYTGG
	2d	WVAQGT	TGGGTNGCNCARGGNAC
	3a	FVLGHD	TTYGTNYTNGGNCAYGA
	3b	VLGHDC	GTNYTNGGNCAYGAYTG
15	3c	GHDCGH	GGNCAYGAYTGYGGNCA
	3d	CGHGSF	TGYGGNCAYGGNWSNTT
	4a	PYHGW	CCNTAYCAYGGNTGG
	4b	HGWRISH	CAYGGNTGGMGNATHWSNCA
	4c-1	WRISHRTHH	TGGMGNATHTCNCAYMGNACNCAYCA*
20	4c-2		TGGMGNATHAGYCAYMGNACNCAYCA*
	4d ·	WRISH	TGGMGNATHWSNCAY
	4e	HRTHH	CAYMGNACNCAYCAY
	5a	ENDESW	GARAAYGAYGARWSNTGG
	5b	DESWVP	GAYGARWSNTGGGTNCC
25	_		
•	6a	WLDAVT	NGTNACNGCRTCNARCCA
•	6b	TYLHH	RTGRTGNARRTANGT
	7a-1	WSYLRGGL	ARNCCNCCNCKNARRTARCTCCA*
0.0	7a-2		ARNCCNCCNCKNARRTANGACCA*
30	7b	LTTIDRD	RTCNCKRTCDATNGTNGTNA
	7c	TIDRDY	RTARTCNCKRTCDATNGT
	8a	HDIGTH	RTGNGTNCCDATRTCRTG
	8b	HVIHHL	NARRTGRTGDATNACRTG
95	8c	HHLFPQI	DATYTGNGGRAANARRTGRTG
35	8d	HLFPQIP	GGDATYTGNGGRAANARRTG
	8e	LFPQIPHY	RTARTGNGGDATYTGNGGRAANA

 $^{^{\}star}$ synthesize 4c and 7a in two pools each to limit the 40 degeneracy

Oligos for 6a - 8e are the complement of the coding sequence

TABLE 8
Table of Oligomers for PCR RACE of fad2

5	Peptide #	Oligo Length	Fold Degeneracy	Similarity with L26296	Similarity in Last 10 n.t.
	la	20	384	75 %	80 %
	1b	17	192	88	80
	1c	17	32	65	80
10			•		
	2a	24	64	79	100
	2b	15	4 8	73	80
	2c	15	48	100	100
	2d	17	128	76	90
15					
	3a	17	. 384	76	70
	3b	17	384	82	80
	3c	17	128	88	90
	3d	17	384	82	70
20		•			
	4a	15	. 64	80	70
	4 b	20	192	75	90
	4c	26	96*	81	80
	4 d	15	216	87	90
25	4e	15	192	87	80
		•			
	5a	18	96	72	80
	5 b	17	96	76	80
00			0.70		
30	6a	18	. 256	78	80
٠	6b	15	192	93	100
	7a	23	256*	78	- 60
	7 b	20	384	90	80
35	7c	18	192	94	90
	8a	18	384	72	70
•	8b	18	192	89	80
	8c	21	384	81	100
40	8d	20	192	80	90 ়
	8e	23	192	83	70
					_

^{*} done in two oligo pools

Table 7 shows the 28 peptide targets from the eight conserved regions and the 30 degenerate oligonucleotides derived from the peptide sequences. The degeneracy was kept to less than 516 fold, for those instances where more degeneracy occurred, by the use of deoxyinosine 5 (Sambrook et al.) and by not including the last nucleotide in the last codon, and in two cases by the use of two subpools. Table 8 shows the amount of degeneracy for each designed oligonucleotide sequence and the amount of homology of the oligonucleotides to the Arabidopsis oleic desaturase fad2 (Accession No. L26296). Also shown in Table 8 is the percent homology in 10 the last 10 nucleotides on the 3' end of each primer, since this region is most important for annealing and elongation under PCR conditions. It is expected that both 10 of 10 and 9 of 10 homology matches, and probably 8 of 10 homology matches in the 3' primer regions will serve as efficient PCR primers. Note that for oligonucleotide sets 1a through 5b (for 3' RACE) the 15 strand direction is the same as the mRNA while for oligonucleotide sets 6a through 8e (for 5' RACE) the direction is opposite of the mRNA. Four oligonucleotides have a 10 of 10 match in the 3' position, 6 oligonucleotides match 9 of 10 in the 3' position and 12 match in 8 of 10 nucleotides in the 3' position. Oligonucleotides corresponding to peptides 2a, 2c, 2d, 3c, 4b, 4d, 20 6b, 7c, 8c, and 8d show 90% or greater homology in their last 10 nucleotides and anneal to the oleic desaturase gene and serve as primers to this gene. This demonstrates the validity of using the conserved regions of the plant linoleic desaturases and DesA to identify and isolate plant oleic desaturases.

The first round of PCR products are subjected to two rounds of subtraction using biotinylated fad3, fadD and fadE cloned cDNA to remove any hybridizing fad3, fadD and fadE sequences with strepavidin. This subtracted DNA is greatly enriched for fad2 sequences and depleted of fad3, fadD and fadE sequences. These 30 samples are run on agarose gels,

blotted and hybridized with pools of probe from the 30 samples. Pools of 5 of each of the 30 PCR samples are labeled with random primers and hybridized to the blots of the 30 samples, for a total of 6 blots hybridized with 6 pools of 5 probes. Additionally, a pool of fad3, fadD and fadE probe is 5 hybridized to a duplicate blot. Bands that do not hybridize strongly to fad3, fadD and fadE but do cross hybridize to probe made from a different sample are strong candidates for fad2 as fad2 is likely to be the only DNA amplified in two or more independent PCR reactions. Positively hybridizing lanes identify samples to amplify by PCR using the same primers as in the initial 10 reaction for 5 to 10 cycles and the PCR products are cloned into plasmid vectors. The same probe that recognized the sample on the blot is used to screen the library and identify the hybridizing clone. Positive clones are sequenced and identified as fad2 clones by their homology but non-identity with fad3, and further characterized as described below.

In the event that fad2 sequences are not sufficiently enriched in one round of PCR to be identified, a second round of PCR is performed. If the lack of detection is due to insufficient amplification of fad2, then another round of PCR using the same primers on the subtracted PCR first round samples and the same simple screen as described above will identify 20 fad2. If there are too many competing non-specific reactions then a second round of PCR using a different primer combination will remove non-specific amplifications and enrich for fad2. To further enrich for fad2 sequences each of the initial 30 PCR samples (one for each oligonucleotide in Table 7) after subtraction as described above, is subjected to a second round of PCR 25 reactions using a different primer combination than the first reaction. One of the primers would be the same degenerate oligonucleotide primer as in the first PCR reaction. The second primer would now be from one of the 30 primers in Table 7 from the opposite class, ie, primers from 1a to 5b form matched sets with primers from 6a to 8e (primers 1a to 5b are in the sense

direction while primers 6a to 8e are in the antisense direction). For example, if oligonucleotide 1a was used initially, it is used again as one of the two primers and the second primer is each of the 6a to 8e oligonucleotides for a total of 11 separate PCR reactions. In total the 30 5 initial reactions result in 418 second cycle PCR reactions, a number easily handled by PCR technology. Essentially this second PCR cycle accomplishes a "nested" or sequential PCR reaction step after removing all the linoleic desaturases by the subtraction step. This increases the amplification as well as the specificity. Identification of samples containing 10 fad2 are performed similarly as described above, with the 418 samples dot blotted onto 22 filters and probed with 21 pools of 20 samples and with a pool of fad3, fadD and fadE. Again, any sample that cross hybridizes with an independent probe sample and does not hybridize to fad3, fadD and fadE is a candidate for containing fad2 in the sample. If fad3, fadD and fadE 15 hybridization is still present, another biotinylation/stepavidin subtraction should remove it. Positively hybridizing samples are run on gels, the band identified by hybridization and isolated for cloning. This second set of PCR reactions produces PCR products of a predictable size since both primers are within the coding region where little variation in size is expected. Thus 20 the presence of a band of the expected size on a gel is diagnostic of fad2, particularly if hybridization of a blot of such a gel with a fad3, fadD and fadE probe indicates the band is not due to fad3, fadD and fadE contamination. After cloning the inserts in E. coli, the resulting plasmids containing the insert are identified by hybridization. They are sequenced 25 and identified as oleic desaturases by their homology but non-identity with the linoleic desaturases, as in the examples described previously. The full length clone of these cDNAs is obtained by standard methods and inserted into plant gene expression and transformation vectors and transformed into Arabidopsis fad2 mutants to confirm the identity of the oleic

desaturase by genetic complemention as was described in the example with linoleic desaturase.

Thus in this approach to isolating the plant oleic desaturases, the total number of peptide regions is 8, comprised of 28 smaller peptide 5 targets. This leads to set of 30 degenerate oligonucleotides, that are used in the PCR amplification and screening of the PCR products. Subtraction of interfering fad3, fadD and fadE sequences is used at several points. If necessary a second round of PCR reactions with paired internal primers gives extra amplification and specificity. This approach identifies the plant 10 oleic desaturases, and the sequence of the isolated clones should confirm their identity by their homology to the plant linoleic desaturases as described. Thus a defined approach to isolating the plant oleic desaturases from the information about linoleic desaturases is presented here. The example given here is for Arabidopsis or canola oleic desaturases, but the 15 approach is not limited to those plants as the oleic desaturases are probably highly conserved in most plants. Thus once one plant oleic desaturase is isolated, the sequence information is used to isolate the genes from other plant species by direct hybridization or by an approach similar to the one described here.

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-69-

SEQUENCE LISTING

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 - (ii) TITLE OF INVENTION: Altered Linolenic and Linoleic Acid Content in Plants
 - (iii) NUMBER OF SEQUENCES: 72
 - (iv) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
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 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25 (EPO)
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 - (A) APPLICATION NUMBER: US 08/014431
 - (B) FILING DATE: 05-FEB-1993
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1353 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 87..1238
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

AATCCATCAA ACCTTTATTC ACCACATTTC ACTGAAAGGC CACACATCTA GAGAGAGAAA 60
CTTCGTCCAA ATCTCTCTCT CCAGCG ATG GTT GTT GCT ATG GAC CAG CGC AGC 113

WO 94/18337 PCT/US94/01321

-70-

Met Val Val Ala Met Asp Gln Arg Ser 1 5

	Val										Glu	GGG Gly				16	1
												GCG Ala				. 20	9
												AGC Ser				25	7
			Phe									GCC Ala 70				30	5
												CAA Gln				35:	3
												CAT His				40:	1
												ATT Ile				449)
												CAT				497	7
												TGG Trp 150				545	;
												CGG Arg				593	3
												TAT Tyr				641	
												TAC Tyr				689)
		Pro					Leu					ACT Thr				737	,
cc	ATA	ATG	TTG	GCC	ACT	CTT	GTT	TAT	CTA	TCG	TTC	CTC	GTT	GAT	CCA	785	;

-71-

Ser	Ile	Met 220	Leu	'Ala	Thr	Leu	Val 225	Tyr	Leu	Ser	Phe	Leu 230	Val	Asp	Pro	٠
						TAT Tyr 240									ATG Met	833
															AAG Lys 265	881
						AAG Lys									TTA Leu	929
						TAC Tyr									•	977
						CAT His										1025
						AGA Arg 320										1073
						TCA Ser										1121
						AAA Lys										1169
						ACA Thr										1217
		TCT Ser 380				TAAC	TTTT	cr 1	(TOOT	AGCTO	T AT	TAGO	ITAA:	A		1265
AACA	CTC	CTT C	TCTI	TTAC	T TA	TTTC	TTTC	TGC	TTT	AGT	TTA	TAAL	TA C	CTCGT	(GAAA)	1325
CTTI	TTTI	TA I	TAAT	GTAI	T	CGTI	AC									1353

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 383 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

- Met Val Val Ala Met Asp Gln Arg Ser Asn Val Asn Gly Asp Ser Gly 1 5 10 15
- Ala Arg Lys Glu Glu Gly Phe Asp Pro Ser Ala Gln Pro Pro Phe Lys
 20 25 30
- Ile Gly Asp Ile Arg Ala Ala Ile Pro Lys His Cys Trp Val Lys Ser 35 40 45
- Pro Leu Arg Ser Met Ser Tyr Val Thr Arg Asp Ile Phe Ala Val Ala 50 55 60
- Ala Leu Ala Met Ala Ala Val Tyr Phe Asp Ser Trp Phe Leu Trp Pro
 65 70 75 80
- Leu Tyr Trp Val Ala Gln Gly Thr Leu Phe Trp Ala Ile Phe Val Leu 85 90 95
- Gly His Asp Cys Gly His Gly Ser Phe Ser Asp Ile Pro Leu Leu Asn 100 105 110
- Ser Val Val Gly His Ile Leu His Ser Phe Ile Leu Val Pro Tyr His 115 120 125
- Gly Trp Arg Ile Ser His Arg Thr His His Gln Asn His Gly His Val 130 135 140
- Glu Asn Asp Glu Ser Trp Val Pro Leu Pro Glu Lys Leu Tyr Lys Asn 145 150 155 160
- Leu Pro His Ser Thr Arg Met Leu Arg Tyr Thr Val Pro Leu Pro Met
 165 170 175
- Leu Ala Tyr Pro Ile Tyr Leu Trp Tyr Arg Ser Pro Gly Lys Glu Gly
 180 185 190
- Ser His Phe Asn Pro Tyr Ser Ser Leu Phe Ala Pro Ser Glu Arg Lys 195 200 205
- Leu Ile Ala Thr Ser Thr Thr Cys Trp Ser Ile Met Leu Ala Thr Leu 210 215 220
- Val Tyr Leu Ser Phe Leu Val Asp Pro Val Thr Val Leu Lys Val Tyr 225 230 235 240
- Gly Val Pro Tyr Ile Ile Phe Val Met Trp Leu Asp Ala Val Thr Tyr
 245 250 255
- Leu His His Gly His Asp Glu Lys Leu Pro Trp Tyr Arg Gly Lys 260 265 270
- Glu Trp Ser Tyr Leu Arg Gly Gly Leu Thr Thr Ile Asp Arg Asp Tyr 275 280 285

-73-

Gly	Ile 290	Phe	Asn	Asn	Ile	His 295	His	увр	Ile	Gly	Thr 300	His	Val	Ile	His		
His 305	Leu	Phe	Pro	Gln	Ile 310	Pro	His	Tyr	His	Leu 315	Val	Авр	Ala	Thr	Arg 320		
Ala	Ala	Lys		Val 325	Leu	Gly	Arg	Tyr	Tyr 330	Arg	Glu	Pro	Lys	Thr 335	Ser		
Gly	Ala	Ile	Pro 340	Ile	His	Leu	Val	Glu 345	Ser	Leu	Val	Ala	Ser 350	Ile	Lys	<i></i>	
Lys	Asp	His 355	Tyr	Val	Ser	Asp	Thr 360	Gly	Авр	Ile	Val	Phe 365	Tyr	Glu	Thr	٠	
Asp	Pro 370	Asp	Leu	Tyr	Val	Tyr 375	Ala	Ser	Авр	Lys	Ser 380	Lys	Ile	Asn			
(2)	INFO	RMAT	rion	FOR	SEQ	ID N	10:3:	:									
	(i)	() (E	A) LE B) TY C) ST	engti (PE : (Rani	HARAC H: 25 nucl DEDNE DGY:	bas leic SS:	e pa acio sino	irs 1									
	(ii)	MOI	ECUI	E T	PE:	CDNA	A										
	(xi)	SEÇ	UENC	E DE	ESCRI	PTIC	on: S	SEQ I	D NO):3:							
GGC	SATGO	TG I	CGGZ	ATGO	SA CO	ATA											2
(2)	INFO	RMAT	NOI	FOR	SEQ	ID N	10:4:										-
	(i)	(A (E) LE 3) TY 3) ST	ngth Pe: Rani	IARAC I: 27 nucl EDNE XGY:	bae eic SS:	e pa acid sinç	irs 1									
	(ii)	MOI	ECUI	E TY	PE:	CDNA	1										
					SCRI			EQ I	D NC	:4:							
	GAGC									•							2
(2)	INFO	RMAT	NOI	FOR	SEQ	ID N	10:5:										
	/ i \	SEC	ITENC	E CH	IARAC	TERI	STIC	:S:									

(A) LENGTH: 21 base pairs(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

	(D) TOPOLOGY: linear			•
	(ii) MOLECULE TYPE: cDNA			
999	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:			
CCG	ATCTCAA GATTACGGAA T		;	21
(2)	INFORMATION FOR SEQ ID NO:6:			
	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 			
	(ii) MOLECULE TYPE: cDNA			
	**			
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:			
TTCC	TTANTGC AGGAGTCGCA TANG			24
(2)	INFORMATION FOR SEQ ID NO:7:			
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 18 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear			
	(ii) MOLECULE TYPE: cDNA			
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:	, .¥:		
AGGA	GTCGCA TAAGGGAG			18
(2)	INFORMATION FOR SEQ ID NO:8:	•		
	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	·		
	(ii) MOLECULE TYPE: cDNA			

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

17

-75-

GGGAAGTGAA	TGGAGAC		

(2) INFORMATION FOR SEQ ID NO:9:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 1645 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(ix) FEATURE:

- (A) NAME/KEY: CDS
- (B) LOCATION: 125..1465

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

·	
GGAAAACACA AGTTTCTCTC ACACACATTA TCTCTTTCTC TATTACCACC ACTCATTCAT	60
AACAGAAACC CACCAAAAAA TAAAAAGAGA GACTTTTCAC TCTGGGGAGA GAGCTCAAGT	120
TCTA ATG GCG AAC TTG GTC TTA TCA GAA TGT GGT ATA CGA CCT CTC CCC Met Ala Asn Leu Val Leu Ser Glu Cys Gly Ile Arg Pro Leu Pro 1 5 10 15	169
AGA ATC TAC ACA ACA CCC AGA TCC AAT TTC CTC TCC AAC AAC AAA Arg Ile Tyr Thr Thr Pro Arg Ser Asn Phe Leu Ser Asn Asn Asn Lys 20 25 30	217
TTC AGA CCA TCA CTT TCT TCT TCT TCT TAC AAA ACA TCA T	265
CTG TCT TTT GGT CTG AAT TCA CGA GAT GGG TTC ACG AGG AAT TGG GCG Leu Ser Phe Gly Leu Asn Ser Arg Asp Gly Phe Thr Arg Asn Trp Ala 50 55 60	313
TTG AAT GTG AGC ACA CCA TTA ACG ACA CCA ATA TTT GAG GAG TCT CCA Leu Asn Val Ser Thr Pro Leu Thr Thr Pro Ile Phe Glu Glu Ser Pro 65 70 75	361
TTG GAG GAA GAT AAT AAA CAG AGA TTC GAT CCA GGT GCG CCT CCT CCG Leu Glu Glu Asp Asn Lys Gln Arg Phe Asp Pro Gly Ala Pro Pro Pro 80 85 90 95	409
TTC AAT TTA GCT GAT ATT AGA GCA GCT ATA CCT AAG CAT TGT TGG GTT Phe Asn Leu Ala Asp Ile Arg Ala Ala Ile Pro Lys His Cys Trp Val 100 105 110	457
AAG AAT CCA TGG AAG TCT TTG AGT TAT GTC GTC AGA GAC GTC GCT ATC Lys Asn Pro Trp Lys Ser Leu Ser Tyr Val Val Arg Asp Val Ala Ile 115 120 125	505

-76-

			GGA Gly						553
			GCT Ala 150					,	601
	 		GGA Gly					·	649
 	 		CAT His						697
			AGT Ser						745
 	 		TCT Ser						793
	•		ACT Thr 230						841
			TTC Phe						889
			CCA Pro						937
			TCT Ser						985
			TTC Phe					1	.033
 	-		TGG Trp 310					1	.081
			GGT Gly					1	.129
			CTG Leu					1	.177

GAC Asp	TAC Tyr	GGA Gly	TTG Leu 355	Ile	AAT Asn	AAC Asn	ATC Ile	CAT His 360	CAT His	GAT Asp	ATT Ile	GGA Gly	ACT Thr 365	CAT His	GTG Val	1225
ATA Ile	CAT Hib	CAT His 370	CTT Leu	TTC Phe	CCG Pro	CAG Gln	ATC Ile 375	CCA Pro	CAT His	TAT Tyr	CAT His	CTA Leu 380	GTA Val	GAA Glu	GCA Ala	1273
Thr	GAA Glu 385	GCA Ala	GCT Ala	AAA Lys	CCA Pro	GTA Val 390	TTA Leu	GGG Gly	AAG Lys	TAT Tyr	TAC Tyr 395	AGG Arg	GAG Glu	CCT Pro	GAT Asp	132:
AAG Lys 400	TCT Ser	GGA Gly	CCG Pro	TTG Leu	CCA Pro 405	TTA Leu	CAT His	TTA Leu	CTG Leu	GAA Glu 410	ATT Ile	CTA Leu	GCG Ala	AAA Lys	AGT Ser 415	1369
ATA Ile	AAA Lys	GAA Glu	GAT Asp	CAT His 420	TAC Tyr	GTG Val	AGC Ser	GAC Asp	GAA Glu 425	GGA Gly	GAA Glu	GTT Val	GTA Val	TAC Tyr 430	TAT Tyr	1417
AAA 1472		GAT	CCA	AAT	CTC	TAT	GGA	GAG	GTC	AAA	GTA	AGA	GCA	GAT	TGAAATGA	AG
		Asp	Pro 435	Asn	Leu	Tyr	Gly	Glu 440	Val	Lys	Val	Arg	Ala 445	Asp		
CAGO	CTT	GAG 2	ATTG/	AAGT	rr rı	CTCT	ATTT(C AG	ACCAC	GCTG	ATT	TTTT	CT !	ract(STATCA	1532
ATTI	TATT	STG :	CAC	CCAC	CA GI	AGAG:	TAG	TA T	CTCT	TAAE	ACG	ATCG	ATC A	AGATO	GAAAC	1592
AAC	AAAT:	rtg :	rttg(CGAT	AC TO	SAAGO	CTAT	A TA	racci	AATA	LAAA	LAAA	AAA i	AAA		164

(2) INFORMATION FOR SEQ ID NO:10:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 446 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

Met Ala Asn Leu Val Leu Ser Glu Cys Gly Ile Arg Pro Leu Pro Arg 1 5 10 15

Ile Tyr Thr Thr Pro Arg Ser Asn Phe Leu Ser Asn Asn Asn Lys Phe 20 25 30

Arg Pro Ser Leu Ser Ser Ser Ser Tyr Lys Thr Ser Ser Ser Pro Leu 35 40 45

Ser Phe Gly Leu Asn Ser Arg Asp Gly Phe Thr Arg Asn Trp Ala Leu 50 55 60

Asn Val Ser Thr Pro Leu Thr Thr Pro Ile Phe Glu Glu Ser Pro Leu

-78-

65					70					75					80
Glu	Glu	Asp	Asn	Lув 85	Gln	Arg	Phe	Авр	Pro 90	Gly	Ala	Pro	Pro	Pro 95	Phe
Asn	Leu	Ala	Asp 100	Ile	Arg	Ala	Ala	Ile 105	Pro	Lys	His	Сув	Trp 110	Val	Lys
Asn	Pro	Trp 115	Lys	Ser	Leu	Ser	Tyr 120	Val	Val	Arg	Авр	Val 125	Ala	Ile	Val
Phe	Ala 130	Leu	Ala	Ala	Gly	Ala 135	Ala	Tyr	Leu	Asn	Asn 140	Trp	Ile	Val	Trp
Pro 145	Leu	Tyr	Trp	Leu	Ala 150	Gln	Gly	Thr	Met	Phe 155	Trp	Ala	Leu	Phe	Val 160
Leu	Gly	His	Авр	Сув 165	Gly	His	Gly	Ser	Phe 170	Ser	Asn	Авр	Pro	Lys 175	Leu
Asn	Ser	Val	Val 180	Gly	His	Leu	Leu	His 185	Ser.	Ser	Ile	Leu	Val 190	Pro	Tyr
His	Gly	Trp 195	Arg	Ile	Ser	His	Arg 200	Thr	His	His	Gln	Asn 205	His	Gly	His
Val	Glu 210	Asn	Asp	Glu	Ser	Trp 215	His	Pro	Met	Ser	Glu 220	Lys	Ile	Tyr	Asn
Thr 225	Leu	Asp	Lys	Pro	Thr 230	Arg	Phe	Phe	Arg	Phe 235	Thr	Leu	Pro	Leu	Val 240
Met	Leu	Ala	Tyr	Pro 245	Phe	Tyr	Leu	Trp	Ala 250	Arg	Ser	Pro	Gly	Lys 255	Lys
Gly	Ser	His	Tyr 260	His	Pro	Авр	Ser	Asp 265	Leu	Phe	Leu	Pro	Lys 270	Glu	Arg
Lys	Asp	Val 275		Thr	Ser	Thr	Ala 280		Trp	Thr	Ala	Met 285	Ala	Ala	Leu
Leu	Val 290	Cys	Leu	Asn	Phe	Thr 295	Ile	Gly	Pro	Ile	Gln 300	Met	Leu	Lys	Leu
Tyr 305	Gly	Ile	Pro	Tyr	Trp 310	Ile	Asn	Val	Met	Trp 315	Leu	Авр	Phe	Val	Thr 320
Tyr	Leu	His	His	His 325	Gly	His	Glu	Asp	Lys 330	Leu	Pro	Trp	Tyr	Arg 335	Gly
Lys	Glu	Trp	Ser 340	Tyr	Leu	Arg	Gly	Gly 345	Leu	Thr	Thr	Leu	Авр 350	Arg	Asp
Tyr	Gly	Leu 355	Ile	Asn	Asn	Ile	His 360	His	Asp	Ile	Gly	Thr 365	His	Val	Ile

WO 94/18337 PCT/US94/01321

-79-

His	His 370	Leu	Phe	Pro	Gln	Ile 375	Pro	His	Tyr	His	Leu 380	Val	Glu	Ala	Thr
Glu 385	Ala	Ala	Lys	Pro	Val 390	Leu	Gly	Lys	Tyr	Tyr 395	Arg	Glu	Pro	Asp	Lys 400
Ser	Gly	Pro	Leu	Pro 405	Leu	His	Leu	Leu	Glu 410	Ile	Leu	Ala	Lys	Ser 415	Ile
Lys	Glu	Asp	His 420	Tyr	Val	Ser	Авр	Glu 425	Gly	Glu	Val	Val	Tyr 430	Tyr	Lys
Ala	Авр	Pro 435	Asn	Leu	Tyr	Gly	Glu 440	Val	Lys	Val	Arg	Ala 445	Asp		

(2) INFORMATION FOR SEQ ID NO:11:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 1525 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(ix) FEATURE:

- (A) NAME/KEY: CDS
- (B) LOCATION: 61..1368

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

AGA	GAGT	GCA 2	ATA	GAAC	GA C	AGAG	ACTT:	r TT	CCTC	TTTT	CTT	CTTG	GGA :	AGAG	CTCC	Α .	60
														CCC Pro 15			108
		–								_				CCC Pro			156
														AAT Asn			204
														GTG Val			252
														GAG Glu			300

-80-

						CCT Pro											348
						TGG											396
			•			GCT Ala										÷	444
						CTT Leu 135											492
						CTC Leu											540
						CCG Pro											588
Leu	His	Ser	Ser 180	Ile	Leu	GTC Val	Pro	Tyr 185	His	Gly	Trp	Arg	Ile 190	Ser	His		636
Arg	Thr	His 195	His	Gln	Asn	CAT	Gly 200	His	Val	Glu	Asn	Авр 205	Glu	Ser	Trp		684
His	Pro 210	Leu	Pro	Glu	Ser	ATC Ile 215	Tyr	Lys	Asn	Leu	Glu 220	Lys	Thr	Thr	Gln		732
Met 225	Phe	Arg	Phe	Thr	Leu 230	Pro	Phe	Pro	Met	Leu 235	Ala	Tyr	Pro	Phe	Tyr 240		780
Leu	Trp	Asn	Arg	Ser 245	Pro	GGG	Lys	Gln	Gly 250	Ser	His	Tyr	His	Pro 255	Asp		828
Ser	Авр	Leu	Phe 260	Leu	Pro	AAA Lys	Glu	Lys 265	Lys	Asp	Val	Leu	Thr 270	Ser	Thr		876
						GCT Ala											924
						CTC Leu 295											972

-81-

TTT Phe 305	GTA Val	ATG Met	TGG Trp	TTG Leu	GAC Asp 310	TTC Phe	GTC Val	ACT Thr	TAC Tyr	TTG Leu 315	CAC His	CAC His	CAT His	GGA Gly	CAT His 320	1020
GAA Glu	GAC Asp	AAG Lys	CTC Leu	CCT Pro 325	TGG Trp	TAT Tyr	CGT Arg	GGA Gly	AAG Lys 330	GAA Glu	TGG Trp	AGT Ser	TAC Tyr	CTG Leu 335	AGA Arg	1068
GGA Gly	GGG Gly	CTC Leu	ACA Thr 340	ACA Thr	TTA Leu	GAT Asp	CGT Arg	GAC Asp 345	TAC Tyr	GGA Gly	TGG Trp	ATC Ile	AAT Asn 350	AAC Aen	ATC Ile	1116
CAC His	CAC His	GAT Asp 355	ATT Ile	GGA Gly	ACT Thr	CAT His	GTG Val 360	ATA Ile	CAT His	CAT His	CTT Leu	TTC Phe 365	CCG Pro	CAG Gln	ATC. Ile	1164
CCA Pro	CAT His 370	TAT Tyr	CAT His	CTA Leu	GTA Val	GAA Glu 375	GCA Ala	ACA Thr	GAA Glu	GCA Ala	GCT Ala 380	AAA Lys	CCA Pro	GTA Val	CTA Leu	1212
GGA Gly 385	AAG Lys	TAC Tyr	TAC Tyr	AGA Arg	GAA Glu 390	CCG Pro	AAA Lys	AAC Asn	TCT Ser	GGA Gly 395	CCT Pro	CTG Leu	CCA Pro	CTT Leu	CAC His 400	1260
TTA Leu	CTG Leu	GGA Gly	AGC Ser	CTC Leu 405	ATA Ile	AAG Lys	AGT Ser	ATG Met	AAA Lys 410	CAA Gln	GAC Asp	CAT His	TTC Phe	GTA Val 415	AGC Ser	1308
			GAT Asp 420													1356
	AGA Arg		TGAG	GACI	ATA C	CTGC/	AGTG!	AA C	CAGG	CAGAC	C AAC	STTAC	ATA			1405
AATI	CATO	CTT C	GCCC	CATTO	A TI	TATG	TCT:	TTI	rgtt:	PTGG	TGT	AAAG	CT 1	TTC	AGATT	1465
LAAA	AAAG	CAT 1	TAA1	TGT	AG AA	AACC:	rgtg(TA	· AAAC:	CTC	GATO	CAAAT	rga i	LATA	GATAT	1525

(2) INFORMATION FOR SEQ ID NO:12:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 435 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

Met Ala Ser Ser Val Leu Ser Glu Cys Gly Phe Arg Pro Leu Pro Arg

									-82-	•					
Phe	Tyr	Pro	Lys 20	His	Thr	Thr	Ser	Phe 25	Ala	Ser	Asn	Pro	Lys 30	Pro	Thi
Phe	Lув	Phe 35	Asn	Pro	Pro	Leu	Lys 40	Pro	Pro	Ser	Ser	Leu 45	Leu	Asn	Se
Arg	Tyr 50	Gly	Phe	Tyr	Ser	Lys 55	Thr	Arg	Asn	Trp	Ala 60	Leu	Asn	Val	Ala
Thr 65	Pro	Leu	Thr	Thr	Leu 70	Gln	Ser	Pro	Ser	Glu 75	Glu	Авр	Thr	Glu	Arq 80
Phe	Asp	Pro	Gly	Ala 85	Pro	Pro	Pro	Phe	Авл 90	Leu	Ala	Авр	Ile	Arg 95	Ala
Ala	Ile	Pro	Lys 100	His	Сув	Trp	Val	Lys 105	Asn	Pro	Trp	Met	Ser 110	Met	Sei
Tyr	Val	Val 115	Arg	Asp	Val	Ala	Ile 120	Val	Phe	Gly	Leu	Ala 125	Ala	Val	Ala
Ala	Tyr 130	Phe	Asn	Asn	Trp	Leu 135	Leu	Trp	Pro	Leu	Tyr 140	Trp	Phe	Ala	Glr
Gly 145	Thr	Met	Phe	Trp	Ala 150	Leu	Phe	Val	Leu	Gly 155	His	Asp	Сув	Gly	Hi:
Gly	Ser	Phe	Ser	Asn 165	Asp	Pro.	Arg	Leu	Asn 170	Ser	Val	Ala	Gly	His 175	Leu
Leu	His		Ser 180	Ile	Leu	Val	Pro	Tyr 185	His	Gly	Trp	Arg	Ile 190	Ser	Hie
Arg	Thr	His 195	His	Gln	Asn	His	Gly 200	His	Val	Glu	Asn	Авр 205	Glu	Ser	Tri
His	210					215					220				
Met 225	Phe	Arg	Phe	Thr	Leu 230	Pro	Phe	Pro	Met	Leu 235	Ala	Tyr	Pro	Phe	Tyr 240

Leu Trp Asn Arg Ser Pro Gly Lys Gln Gly Ser His Tyr His Pro Asp 245

Ser Asp Leu Phe Leu Pro Lys Glu Lys Lys Asp Val Leu Thr Ser Thr

Ala Cys Trp Thr Ala Met Ala Ala Leu Leu Val Cys Leu Asn Phe Val

Met Gly Pro Ile Gln Met Leu Lys Leu Tyr Gly Ile Pro Tyr Trp Ile 290 295

Phe Val Met Trp Leu Asp Phe Val Thr Tyr Leu His His Gly His 305 310

-83-

Glu Asp Lys Leu Pro Trp Tyr Arg Gly Lys Glu Trp Ser Tyr Leu Arg 325 330 335

Gly Gly Leu Thr Thr Leu Asp Arg Asp Tyr Gly Trp Ile Asn Asn Ile 340 345 350

His His Asp Ile Gly Thr His Val Ile His His Leu Phe Pro Gln Ile 355 360 365

Pro His Tyr His Leu Val Glu Ala Thr Glu Ala Ala Lys Pro Val Leu 370 375 380

Gly Lys Tyr Tyr Arg Glu Pro Lys Asn Ser Gly Pro Leu Pro Leu His 385 390 395 400

Leu Leu Gly Ser Leu Ile Lys Ser Met Lys Gln Asp His Phe Val Ser 405 410 415

Asp Thr Gly Asp Val Val Tyr Tyr Glu Ala Asp Pro Lys Leu Asn Gly 420 425 430

Gln Arg Thr 435

- (2) INFORMATION FOR SEQ ID NO:13:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (synthetic)
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:

GAYATHMGNG CNGCNATHCC

20

- (2) INFORMATION FOR SEQ ID NO:14:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 17 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (synthetic)
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

GCNATHCCNA ARCAYTG

17

(2)	INFORMATION FOR SEQ ID NO:15:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 17 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (synthetic)	·
•		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:	
AAR	CAYTGYT GGGTNAA	1
(2)	INFORMATION FOR SEQ ID NO:16:	
	AL ADAMAN MIND AND TONICO	•
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (synthetic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:	
TGG:	ITYYTNT GGCCNYTNTA YTGG	2
(2)	INFORMATION FOR SEQ ID NO:17:	
•	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 15 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	·
	(ii) MOLECULE TYPE: DNA (synthetic)	•
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:	
mcc:		
1667	TTYYTNT GGCCN	1
(2)	INFORMATION FOR SEQ ID NO:18:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 15 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: Bingle	

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (synthetic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:	
(XI) BEGODINGS SECONDICTION DEE IS NOTED	
TGGCCNYTNT AYTGG	15
(2) INFORMATION FOR SEQ ID NO:19:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 17 base pairs	
(B) TYPE: nucleic acid	
(C) STRANDEDNESS: single	
(D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (synthetic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:	
TGGGTNGCNC ARGGNAC	17
(2) INFORMATION FOR SEQ ID NO:20:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 17 base pairs	
(B) TYPE: nucleic acid	•
(C) STRANDEDNESS: single	
(D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (synthetic)	
•	
A LA CRESTANCE PROGRAMMENT AND TRANSPORT	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:	
TTYGTNYTNG GNCAYGA	17
(2) INFORMATION FOR SEQ ID NO:21:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 17 base pairs	
(B) TYPE: nucleic acid	
(C) STRANDEDNESS: Bingle	
(D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (synthetic)	
. *	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:	
GTNYTNGGNC AYGAYTG	

-86-

(4)) INFORMATION FOR SEQ ID NO.22:	
	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	(ii) MOLECULE TYPE: DNA (synthetic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:	•
GGN	NCAYGAYT GYGGNCA	1
(2)	INFORMATION FOR SEQ ID NO:23:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (synthetic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:	
TGY	GGNCAYG GNWSNTT	1
(2)	INFORMATION FOR SEQ ID NO:24:	
	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 15 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	(ii) MOLECULE TYPE: DNA (synthetic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:	
CCN'	TAYCAYG GNTGG	1
(2)	INFORMATION FOR SEQ ID NO:25:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (synthetic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:25	:
CAYGGNTGGM GNATHWSNCA	20
(2) INFORMATION FOR SEQ ID NO:26:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs	
(B) TYPE: nucleic acid	
(C) STRANDEDNESS: single	·
(D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (synthetic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:26	:
TGGMGNATHT CNCAYMGNAC NCAYCA	26
TODIONATHI CHOATHONAC NOATOA	. 20
(2) INFORMATION FOR SEQ ID NO:27:	•
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 26 base pairs	·
(B) TYPE: nucleic acid (C) STRANDEDNESS: single	
(D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (synthetic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:	•
TGGMGNATHA GYCAYMGNAC NCAYCA	26
(2) INFORMATION FOR SEQ ID NO:28:	•
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 15 base pairs	
(B) TYPE: nucleic acid (C) STRANDEDNESS: single	
(C) STRANDEDNESS: Bingle (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (synthetic)	•
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:	
(XI) SEQUENCE DESCRIPTION: SEQ ID NO:28:	•
TOO WON'S MITTE ON ON Y	10

(2)	INFORMATION FOR SEQ ID NO:29:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 15 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
•	(D) TOPOLOGY: linear	
	(2) 101010111	
	(ii) MOLECULE TYPE: DNA (synthetic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:	
CAY	MGNACNC AYCAY	1
(2)	INFORMATION FOR SEQ ID NO:30:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 18 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(b) 10102001. 12	
	(ii) MOLECULE TYPE: DNA (synthetic)	
GARI	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:	1
(2)	INFORMATION FOR SEQ ID NO:31:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 17 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (synthetic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:	·
GAY	GARWSNT GGGTNCC	1
(2)	INFORMATION FOR SEQ ID NO:32:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 18 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: Bingle	
	(D) TOPOLOGY: linear	

(ii) MOLECULE TYPE: DNA (synthetic)	·	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:		
NGTNACNGCR TCNARCCA		18
(2) INFORMATION FOR SEQ ID NO:33:		•
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 15 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 		
(ii) MOLECULE TYPE: DNA (synthetic)	×	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:		
RTGRTGNARR TANGT		15
(2) INFORMATION FOR SEQ ID NO:34:		
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 23 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 		
(ii) MOLECULE TYPE: DNA (synthetic)		
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:		
ARNCCNCCNC KNARRTARCT CCA		23
(2) INFORMATION FOR SEQ ID NO:35:		
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 23 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 		
(ii) MOLECULE TYPE: DNA (synthetic)		
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:		
ARNCCNCCNC KNARRTANGA CCA		23

(2) INFORMATION FOR SEQ ID NO:36:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(5) 2020021 22.1502	:
	(ii) MOLECULE TYPE: DNA (synthetic)	
		·
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:	
2000		_
RTC	CNCKRTCD ATNGTNGTNA	2
(2)) INFORMATION FOR SEQ ID NO:37:	
` .	,	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 18 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (synthetic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:	
RTA	ARTCNCKR TCDATNGT	18
(2)	INFORMATION FOR SEQ ID NO:38:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 18 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (synthetic)	
		•
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:	
RTG	NGTNCCD ATRICRIG	18
(2)	INFORMATION FOR SEQ ID NO:39:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 18 base pairs	

(B) TYPE: nucleic acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (synthetic)			•
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:			
NARRIGRIGD ATNACRIG			18
(2) INFORMATION FOR SEQ ID NO:40:	•	·	•
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	·		
(ii) MOLECULE TYPE: DNA (synthetic)			
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:			
DATYTGNGGR AANARRTGRT G			21
(2) INFORMATION FOR SEQ ID NO:41:			
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (synthetic)			
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:			
GGDATYTGNG GRAANARRTG			20
(2) INFORMATION FOR SEQ ID NO:42:	•		•
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 23 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 			
(ii) MOLECULE TYPE: DNA (synthetic)			
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:42:			
RTARTGNGGD ATYTGNGGRA ANA			23

- (2) INFORMATION FOR SEQ ID NO:43:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 7 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:43:

Asp Ile Arg Ala Ala Ile Pro

- (2) INFORMATION FOR SEQ ID NO:44:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:44:

Ala Ile Pro Lys His Cys
1 5

- (2) INFORMATION FOR SEQ ID NO:45:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:45:

Lys His Cys Trp Val Lys
1 5

- (2) INFORMATION FOR SEQ ID NO:46:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 8 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:46:

Trp Phe Leu Trp Pro Leu Tyr Trp 1

- (2) INFORMATION FOR SEQ ID NO:47:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:47:

Trp Phe Leu Trp Pro

- (2) INFORMATION FOR SEQ ID NO:48:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:48:

Trp Pro Leu Tyr Trp
1 5

- (2) INFORMATION FOR SEQ ID NO:49:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:49:

Trp Val Ala Gln Gly Thr

(2) INFORMATION FOR SEQ ID NO:50:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:50:

Trp Val Ala Gln Gly Thr

- (2) INFORMATION FOR SEQ ID NO:51:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:51:

Val Leu Gly His Asp Cys 1 5

- (2) INFORMATION FOR SEQ ID NO:52:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:52:

Gly His Asp Cys Gly His 1 5

- (2) INFORMATION FOR SEQ ID NO:53:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide

-95-

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:53:

Cys Gly His Gly Ser Phe
1 5

- (2) INFORMATION FOR SEQ ID NO:54:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:54:

Pro Tyr His Gly Trp 1 5

- (2) INFORMATION FOR SEQ ID NO:55:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 7 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:55:

His Gly Trp Arg Ile Ser His

- (2) INFORMATION FOR SEQ ID NO:56:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 9 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:56:

Trp Arg Ile Ser His Arg Thr His His 1

- (2) INFORMATION FOR SEQ ID NO:57:
 - (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 5 amino acids
- (B) TYPE: amino acid
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:57:

Trp Arg Ile Ser His

- (2) INFORMATION FOR SEQ ID NO:58:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:58:

His Arg Thr His His 1 5

- (2) INFORMATION FOR SEQ ID NO:59:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:59:

Glu Asn Asp Glu Ser Trp 1 5

- (2) INFORMATION FOR SEQ ID NO:60:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide

-97-

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:60:

Asp Glu Ser Trp Val Pro 1 5

- (2) INFORMATION FOR SEQ ID NO:61:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:61:

Trp Leu Asp Ala Val Thr

- (2) INFORMATION FOR SEQ ID NO:62:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:62:

Thr Tyr Leu His His 1 5

- (2) INFORMATION FOR SEQ ID NO:63:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 8 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:63:

Trp Ser Tyr Leu Arg Gly Gly Leu

- (2) INFORMATION FOR SEQ ID NO:64:
 - (i) SEQUENCE CHARACTERISTICS:

-98-

- (A) LENGTH: 7 amino acids
- (B) TYPE: amino acid
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:64:

Leu Thr Thr Ile Asp Arg Asp
1 5

- (2) INFORMATION FOR SEQ ID NO:65:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:65:

Thr Ile Asp Arg Asp Tyr
1 5

- (2) INFORMATION FOR SEQ ID NO:66:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:66:

His Asp Ile Gly Thr His

- (2) INFORMATION FOR SEQ ID NO:67:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide

-99-

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:67:

His Val Ile His His Leu 1 5

- (2) INFORMATION FOR SEQ ID NO:68:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 7 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:

His His Leu Phe Pro Gln Ile
1 5

- (2) INFORMATION FOR SEQ ID NO:69:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 7 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:69:

His Leu Phe Pro Gln Ile Pro 1 5

- (2) INFORMATION FOR SEQ ID NO:70:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: B amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:70:

Leu Phe Pro Gln Ile Pro His Tyr

- (2) INFORMATION FOR SEQ ID NO:71:
 - (i) SEQUENCE CHARACTERISTICS:

-100-

(A) LENGTH: 1670 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 46..1302

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:71:

	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:71:															
CAAACTCTCT CGGGGGGTCG CTTCTTCTGC ATTTTCTGCT TC														CT TO la Se		54
														TGT Cys		102
														CCT Pro		150
					Pro									ACT Thr 50		198
														TGT Cys		246
														GCA Ala		294
														GGA Gly		342
														CTT Leu		390
														GTG Val 130		438
ATA Ile	TCT Ser	GTG Val	ACT Thr 135	TCA Ser	TAC Tyr	ACT	TTG Leu	GGG Gly 140	CTC Leu	TTC Phe	ATG Met	ATT Ile	GCA Ala 145	AAA Lys	TCG Ser	486
CCG	TGG	TAT	CTG	CTA	CCG	TTG	GCT	TGG	GCA	TGG	ACA	GGA	ACT	GCA	ATT	534

Pro	Trp	Tyr 150	Leu	Leu	Pro	Leu	Ala 155	Trp	Ala	Trp	Thr	Gly 160	Thr	Ala	Ile	
												AAG Lys				582
	Asn											GCC Ala				630
												GAC Asp				678
												CAG Gln				726
												GCA Ala 240				774
												CÁC His				822
												GTG Val				870
												GTT Val				918
												AAA Lys				966
ATG Met	CCA Pro	TGG Trp 310	TTG Leu	GGC	TAT Tyr	CAC His	TTC Phe 315	TGG Trp	ATG Met	AGC Ser	ACA Thr	TTC Phe 320	ACA Thr	ATG Met	GTT Val	1014
												GAT Asp				1062
GCG Ala 340	GCT Ala	CAG Gln	GCC Ala	CAG Gln	CTG Leu 345	AAT Asn	GGA Gly	ACT Thr	GTT Val	CAT His 350	TGT Cyb	GAC Asp	TAC Tyr	CCT Pro	AGT Ser 355	1110
TGG Trp	ATT Ile	GAA Glu	ATT Ile	CTC Leu 360	TGC Cyb	CAT His	GAT Asp	ATC Ile	AAC Asn 365	GTT Val	CAC His	ATC Ile	CCG Pro	CAT His 370	CAT His	1158
ATT	AGC	CCA	AGA	ATA	CCG	AGC	TAC	AAT	CTC	CGT	GCA	GCT	CAT	GAG	TCT	1206

1669

1670

-102-

Ile	Ser	Pro	Arg 375		Pro	Ser	Tyr	Asn 380	Leu	Arg	Ala	Ala	His 385	Glu	Ser		
														AAC			125
Ile	Gln	390		Trp	Gly	Lув	Tyr 395	Thr	Asn	Leu	Ala	Thr 400	Trp	Asn	Trp		
CGA 1309		ATG	AAG	ACG	ATA	ATG	ACT	GTG	TGT	CAT	GTC	TAT	GAC	AAA	TAGG	AGAA	T
Arg	Leu 405		Lys	Thr	Ile	Met 410	Thr	Val	Сув	His	Val 415	Tyr	Авр	Lys		·	
ACAT	TCC	TT :	TGACC	CGGTI	TA GO	ccci	GAAG	CAA :	CTC	GCC	ATAA	ACCI	TC (CTCAP	GAAA!	r	1369
CAAI	GCC	AA (CTACA	CAGO	C TG	ATTO	GCCA	TGG	TCTC	AAA	CTAG	TCTI	TT (GAAAI	CTCA	A .	1429
TATO	TTT	TG (CAGTO	GCCG	A TG	TTAT	ATGI	' AAG	CTTI	CCA	AGCG	ATGA	GC :	TTCTC	TAAC	A	1489
CTTC	ACCA	LAC (CTTI	CATAC	T GI	TATO	TTCI	TTC	CAAT	CTT	ATCA	GAAG	AG I	AGAAA	CTGG	r	1549
CAAA	IATT.	CT (GAGCG	ATTG	C AA	TTCI	'TTTA	TCA	GTTI	CTT	AGCT	'ATAA	GA I	AGATT	GAAC	A	1609

(2) INFORMATION FOR SEQ ID NO:72:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 418 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:72:

Met Ala Ser Arg Ile Ala Asp Ser Leu Phe Ala Phe Thr Gly Pro Gln
1 5 10 15

Gln Cys Leu Pro Arg Val Pro Lys Leu Ala Ala Ser Ser Ala Arg Val 20 25 30

Ser Pro Gly Val Tyr Ala Val Lys Pro Ile Asp Leu Leu Lys Gly
35 40 45

Arg Thr His Arg Ser Arg Arg Cys Val Ala Pro Val Lys Arg Arg Ile 50 55 60

Gly Cys Ile Lys Ala Val Ala Ala Pro Val Ala Pro Pro Ser Ala Asp
65 70 75 80

Ser Ala Glu Asp Arg Glu Gln Leu Ala Glu Ser Tyr Gly Phe Arg Gln 85 90 95

Ile	Gly	Glu	Asp 100	Leu	Pro	Glu	Asn	Val 105	Thr	Leu	Lys	Авр	11e 110	Met	Авр
Thr	Leu	Pro 115	Lys	Glu	Val	Phe	Glu 120	Ile	Авр	Asp	Leu	Lув 125	Ala	Leu	Lys
Ser	Val 130	Leu	Ile	Ser	Val	Thr 135	Ser	Tyr	Thr	Leu	Gly 140	Leu	Phe	Met	Ile
145			Pro		150					155					160
			Thr	165					170	٠				175	
			Lys 180					185					190		
		195	Leu		-		200					205			
	210		Ala			215					220				
225			Pro		230					235					240
			Gly	245					250					255	
_	· ,		Trp 260					265					270		
		275	Lys		•		280					285			
	290		Leu Met			295					300				
305	_		His		310					315				•	320
			Ala	325		,			330		٠			335	
			340 Trp					345					350		
-		355	Ile				360					365			
	370		ile			375					380				
385	GIU	net		11	390		1	,	_, _	395					400

-104-

Trp Asn Trp Arg Leu Met Lys Thr Ile Met Thr Val Cys His Val Tyr
405 410 415

Asp Lys

10

Claims:

- 1. A genetically transformed plant which has an elevated linolenic acid content comprising a recombinant, double-stranded DNA molecule comprising
 - (i) a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;
 - (ii) a structural coding sequence that causes the production of an RNA sequence that encodes a linoleic acid desaturase activity; and
 - (iii) a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence.
- 15 2. The plant of claim 1 in which the linoleic acid desaturase activity is from plants.
 - 3. The plant of claim 1 in which the linoleic acid desaturase activity is from fungi, algae or bacteria.
- 4. The plant of claim 1 in which the structural coding 20 sequence of (ii) is taken from SEQ. ID NO:1.
 - 5. The plant of claim 1 in which the structural coding sequence of (ii) is taken from SEQ. ID NO:9.
 - 6. The plant of claim 1 in which the structural coding sequence of (ii) is taken from SEQ. ID NO:11.
- 7. The plant of claim 1 in which the promoter of (i) is an endogenous plant linoleic acid desaturase promoter.
 - 8. A genetically transformed plant which has a reduced linolenic acid content, comprising a recombinant, double-stranded DNA molecule comprising

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- (i) a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;
- (ii) a DNA sequence that causes the production of an RNA sequence that is in antisense orientation to at least a portion of a gene that encodes a linoleic acid desaturase activity in said plant; and
- (iii) a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence.
- 9. The plant of claim 8 in which the linoleic acid desaturase enzyme is from plants.
- 10. The plant of claim 8 in which the linoleic acid desaturase enzyme is from fungi, algae or bacteria.
- 15 11. The plant of claim 8 in which the structural coding sequence of (ii) is taken from SEQ. ID NO:1.
 - 12. The plant of claim 8 in which the structural coding sequence of (ii) is taken from SEQ. ID NO:9.
- 13. The plant of claim 8 in which the structural coding 20 sequence of (ii) is taken from SEQ. 8 ID NO:11.
 - 14. The plant of claim 8 in which the promoter of (i) is an endogenous plant linoleic acid desaturase promoter.
- 15. A genetically transformed plant which has an improved resistance to low temperatures comprising a recombinant, double-stranded
 25 DNA molecule comprising
 - (i) a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;

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(ii)	а	structural	coding	sequence	that	cau	ses	the
produc	tio	n of an RN	A seque	nce that e	ncode	s a	linc	leio
acid de	sat	urase activ	ity; and					

- (iii) a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence.
- 16. A genetically transformed plant which has an elevated ability to respond to pathogens, comprising a recombinant, double-stranded DNA molecule comprising
 - (i) a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;
 - (ii) a structural coding sequence that causes the production of an RNA sequence that encodes a linoleic acid desaturase activity; and
 - (iii) a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence.
- 17. A seed produced from genetically transformed plant where 20 said seed has an linolenic acid content suitable for use as a source of linolenic acid, said plant comprising a recombinant, double-stranded DNA molecule comprising
 - (i) a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;
 - (ii) a structural coding sequence that causes the production of an RNA sequence that encodes a linoleic acid desaturase activity; and

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- (iii) a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence.
- 18. The seed of claim 17 where said plant is selected from the 5 group consisting of soybean and rapeseed.
 - 19. A genetically transformed plant which has a linolenic acid content of less than about 3%, said plant comprising a recombinant, double-stranded DNA molecule comprising
 - (i) a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;
 - (ii) a DNA sequence that causes the production of an RNA sequence that is in antisense orientation to at least a portion of a gene that encodes a linoleic acid desaturase activity in said plant; and
 - (iii) a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence.
- 20. A genetically transformed plant which has an increased 20 oleic acid content, comprising a recombinant, double-stranded DNA molecule comprising
 - (i) a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;
 - (ii) a DNA sequence that causes the production of an RNA sequence that is in antisense orientation to at least a portion of a gene that encodes a oleic acid desaturase activity in said plant; and

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- (iii) a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence.
- 21. A genetically transformed plant which has an increased 5 oleic acid content, comprising a recombinant, double-stranded DNA molecule comprising
 - (i) a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;
 - (ii) a DNA sequence that causes the production of an RNA sequence that is in antisense orientation to at least a portion of a gene that encodes a linoleic acid desaturase activity in said plant; and
 - (iii) a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence.
 - 22. A method of producing a genetically transformed plant which has an elevated linolenic acid content, comprising
 - inserting into the genome of a plant cell a (a) recombinant, double-stranded DNA molecule comprising:
 - a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;
 - a structural coding sequence that causes the production of an RNA sequence that encodes a linoleic acid desaturase activity; and
 - a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence;

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- (b) obtaining transformed plant cells; and
- (c) regenerating from the transformed plant cells genetically transformed plants which have an elevated linolenic acid content.
- 5 23. The method of claim 22 in which the linoleic acid desaturase enzyme is from plants.
 - 24. The method of claim 22 in which the linoleic acid desaturase enzyme is from fungi, algae or bacteria.
- 25. The method of claim 22 in which the structural coding 10 sequence of (ii) is taken from SEQ. ID NO:1.
 - 26. The method of claim 22 in which the structural coding sequence of (ii) is taken from SEQ. ID NO:9.
 - 27. The method of claim 22 in which the structural coding sequence of (ii) is taken from SEQ. ID NO:11.
- 15 28. The plant of claim 22 in which the promoter of (i) is an endogenous plant linoleic acid desaturase promoter.
 - 29. A method of producing a genetically transformed plant which has a reduced linolenic acid content, comprising
 - (a) inserting into the genome of a plant cell a recombinant, double-stranded DNA molecule comprising:
 - (i) a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;
 - (ii) a DNA sequence that causes the production of an RNA sequence that is in antisense orientation to at least a portion of a gene that encodes a linoleic acid desaturase activity in said plant; and

- (iii) a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence
- (b) obtaining transformed plant cells; and
- 5 (c) regenerating from the transformed plant cells genetically transformed plants which have a reduced linolenic acid content.
 - 30. The method of claim 29 in which the linoleic acid desaturase enzyme is from plants.
- 10 31. The method of claim 29 in which the linoleic acid desaturase enzyme is from fungi, algae or bacteria.
 - 32. The method of claim 29 in which the structural coding sequence of (ii) is taken from SEQ. ID NO:1.
- 33. The method of claim 29 in which the structural coding sequence of (ii) is taken from SEQ. ID NO:9.
 - 34. The method of claim 29 in which the structural coding sequence of (ii) is taken from SEQ. ID NO:11.
 - 35. The plant of claim 29 in which the promoter of (i) is an endogenous plant linoleic acid desaturase promoter.
- 36. A method of producing a genetically transformed plant which has an increased oleic acid content, comprising
 - (a) inserting into the genome of a plant cell a recombinant, double-stranded DNA molecule comprising:
 - (i) a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;
 - (ii) a DNA sequence that causes the production of an RNA sequence that is in antisense orientation to at least a portion of a

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gene that encodes a linoleic acid desaturase activity in said plant; and

- (iii) a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence
- (b) obtaining transformed plant cells; and
- (c) regenerating from the transformed plant cells genetically transformed plants which have an increased oleic acid content.
- 37. A recombinant, double-stranded DNA molecule comprising in sequence:
 - (i) a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;
 - (ii) a structural coding sequence that causes the production of an RNA sequence that encodes a linoleic acid desaturase activity; and
 - (iii) a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence.
 - 38. A recombinant, double-stranded DNA molecule comprising in sequence:
 - (i) a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;
 - (ii) a DNA sequence that causes the production of an RNA sequence that is in antisense orientation to at least a portion of a gene that encodes a linoleic acid desaturase activity in said plant; and

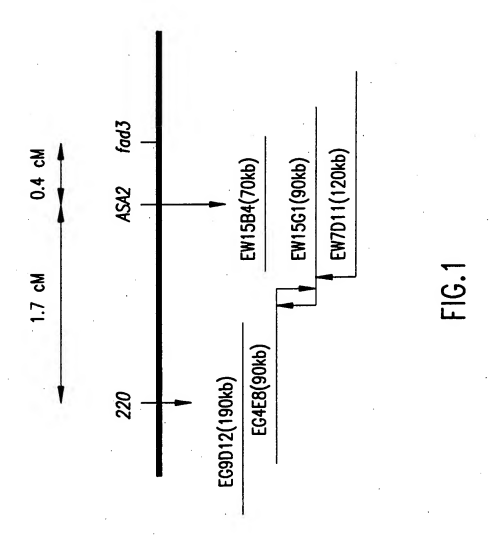
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- (iii) a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence.
- 39. A plant cell comprising a recombinant, double-
- 5 stranded DNA molecule comprising in sequence:
 - (i) a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;
 - (ii) a DNA sequence that causes the production of an RNA sequence that is in antisense orientation to at least a portion of a gene that encodes a linoleic acid desaturase activity in said plant; and
 - (iii) a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence.
 - 40. A method of producing a genetically transformed plant which has an increased oleic acid content, comprising
 - (a) inserting into the genome of a plant cell a recombinant, double-stranded DNA molecule comprising:
 - (i) a promoter that functions in plant cells to cause the production of an RNA sequence, said promoter operably linked to;
 - (ii) a DNA sequence that causes the production of an RNA sequence that is in antisense orientation to at least a portion of a gene that encodes a oleic acid desaturase activity in said plant; and

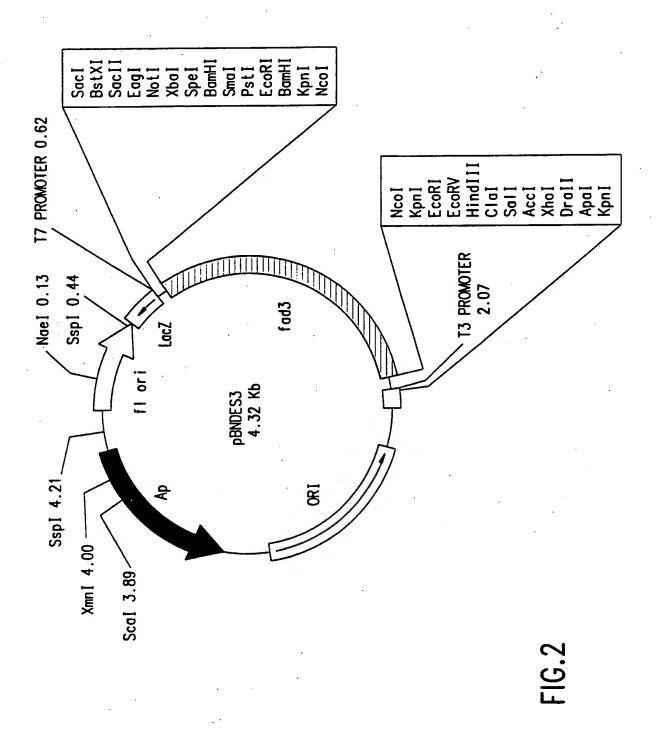
15

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- (iii) a 3' non-translated region that functions in plant cells to promote polyadenylation to the 3' end of said RNA sequence
- (b) obtaining transformed plant cells; and
- (c) regenerating from the transformed plant cells genetically transformed plants which have an increased oleic acid content.



RECTIFIED SHEET (RULE 91)
ISA/EP



RECTIFIED SHEET (RULE 91) ISA/EP

AAT	CCAT	CAA /	ACCT.	TAT	TC A	CCAC	ATTT(C AC	TGAA	AGGC	CAC	ACAT	CTA (GAGA(GAGAAA		60
CTT	CGTC	CAA /	ATCT(CTCT(CT CO	CAGC									C AGC g Ser	1	113
	Vol													GAT Asp	CCA Pro 25	. 1	161
														ATT He 40		2	209
														GTC Vol			257
														TAT Tyr		3	305
														ACC Thr		3	353
														AGT Ser		4	101
														CAT His 120		4	149
														ACA Thr		4	197
								Asn						CCG Pro		5	545

FIG.3a RECTIFIED SHEET (RULE 91) ISA/EP

GAA Glu 155										593
ACT Thr										641
AGT Ser								Leu		689
GCT Alo		Glu								73 7
ATA He										785
ACA Thr 235									•	833
TTG Leu										881
CCT Pro										929
ACT Thr										977
 GGA Gly									,	1025

FIG.3b
RECTIFIED SHEET (RULE 91)
ISA/EP

CAC TIG GIC GAT GCC ACG AGA GCA GCT AAA CAT GIG TTA GGA AGA TAC His Leu Val Asp Ala Thr Arg Ala Ala Lys His Val Leu Gly Arg Tyr 315 320 325	1073
TAC AGA GAG CCG AAG ACG TCA GGA GCA ATA CCG ATT CAC TTG GTG GAG Tyr Arg Glu Pro Lys Thr Ser Gly Ala lle Pro lle His Leu Val Glu 330 345	1121
AGT TTG GTC GCA AGT ATT AAA AAA GAT CAT TAC GTC AGT GAC ACT GGT Ser Leu Vol Alo Ser Ile Lys Lys Asp His Tyr Vol Ser Asp Thr Gly 350 355 360	1169
GAT ATT GTC TTC TAC GAG ACA GAT CCA GAT CTC TAC GTT TAT GCT TCT Asp lie Vol Phe Tyr Glu Thr Asp Pro Asp Leu Tyr Vol Tyr Aig Ser 365 370 375	1217
GAC AAA TCT AAA ATC AAT TAACTTTTCT TCCTAGCTCT ATTAGGAATA Asp Lys Ser Lys IIe Asn 380	1265
AACACTECTT CTCTTTTACT TATTIGTTIC TGCTTTAAGT TTAAAATGTA CTCGTGAAAC	1325
CTTTTTTTA TTAATGTATT TACGTTAC	1353

FIG.3c

Met 1	V a l	Val	Ala	Met 5	Asp	Gln	Arg	Ser	Asn 10	Val	Asn	Gly	Asp	Ser 15	Gly
Ala	Arg	Lys	Glu 20	Glu	Gly	Phe	Asp	Pro 25	Ser	Alo	GIn	Pro	Pro 30	Phe	Lys
He	Gly	Asp 35	He	Arg	Ala	Ala	ile 40	Pro	Ŀys	His	Cyś	Trp 45	Val	Lys	Ser
Pro	Leu 50	Arg	Ser	Met	Ser	Tyr 55	Val	Thr	Arg	Asp	lle 60	Phe	Alo	Val	Alo
Ala 65	Leu	Alo	Met	Alo	A1a 70	Val	Tyr	Phe	Asp	Ser 75	Trp	Phe	Leu	Trp	Pro 80
Leu	Tyr	Trp	Val	A1 a 85	GIn	Gly	Thr	Leu	Phe 90	Trp	Ala	He	Phe	Va 1 95	Leu
Gly	His	Asp	Cys 100	Gly	His	Gly	Ser	Phe 105	Ser	Asp	He	Pro	Leu 110	Leu	Asn
Ser	Val	Va I 115	Gly	His	He	Leu	His 120	Ser	Phe	He	Leu	Vo I 125	Pro	Tyr	His
Gly	Trp 130	Arg	He	Ser	His	Arg 135	Thr	His	His	GIn	Asn 140	His	Gly	His	Vol
G1u 145	Asn	Asp	Glu	Ser	Trp 150	Val	Pro	Leu :	Pro	G I u 155	Lys	Leu	Tyr	Lys	Asn 160
Leu	Pro	His	Ser	Thr 165	Arg	Met	Leu	Arg	Tyr 170	Thr	Val _.	Pro	Leu	Pro 175	Met
Leu	Alo	Tyr	Pro 180	He	Tyr	Leu	Trp	Tyr 185	Arg	Ser	Pro	Gly	Lys 190	Glu	Gly
Ser	His	Phe 195	Asn	Pro	Туг	Ser	Ser 200	Leu	Phe	Ala	Pro	Ser 205	Glu	Arg	Lys
Leu	lle 210	Alo	Thr	Ser	Thr	Thr 215	Cys	Trp	Ser	He	Met 220	Leu	Ala	Thr	Leu
Va I 225	Tyr	Leu	Ser	Phe	Leu 230	Vol	Asp	Pro	Voi	Thr 235	Val	Leu	Lys	Val	Tyr 240
Gly	Val	Pro	Tyr	11e 245	He	Phe	Val	Met	Trp 250	Leu	Asp	Alo	Vol	Thr 255	Tyr

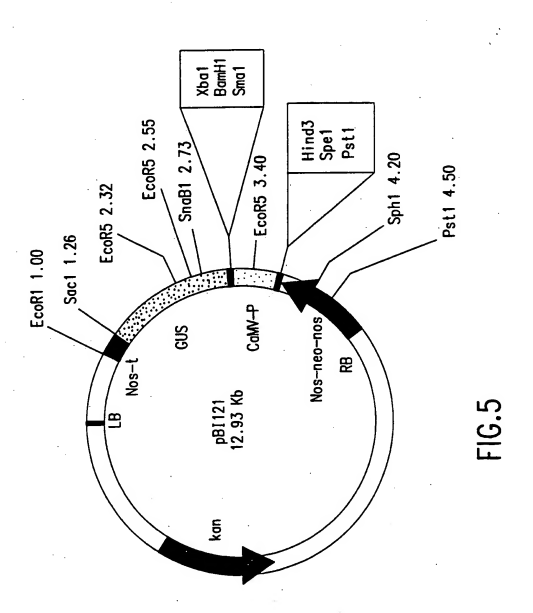
FIG.3d

Leu His His Gly His Asp Glu Lys Leu Pro Trp Tyr Arg Gly Lys Glu Trp Ser Tyr Leu Arg Gly Gly Leu Thr Thr Ile Asp Arg Asp Tyr Gly Ile Phe Asn Asn Ile His His Asp Ile Gly Thr His Val Ile His His Leu Phe Pro Gln Ile Pro His Tyr His Leu Val Asp Ala Thr Arg Alo Alo Lys His Vol Leu Gly Arg Tyr Tyr Arg Glu Pro Lys Thr Ser Gly Alo lle Pro lle His Leu Vol Glu Ser Leu Vol Alo Ser lle Lys Lys Asp His Tyr Val Ser Asp Thr Gly Asp Ile Val Phe Tyr Glu Thr Asp Pro Asp Leu Tyr Val Tyr Ala Ser Asp Lys Ser Lys Ile Asn

FIG.3e

	10				50		
BND3.AMI	RSNVNGDSG/	ARKEEGFDPS/	AQPPFKIGDI	IRAA I PKHCW	VKSPLRSMSY'	VTRDIFAVA. · ·	ALA
DESA.AMI	10				-	VLITLGAIA 50 120	VGY 60
BND3.AMI		LWPLYWVAQ			SDIPLLNSVV		VPY
DESA.AMI		CLPITWIWTO	B0	90	AKKRWVNDLV 100	110	:. YPF
BND3.AM1	130 HGWRISHRTH			160 KLYKNLPHST	170 RMLRYTVPLPI	180 H-LAYPIYL'	WYR :
DESA.AMI	120 13	30 14	40 '	150		170	W
BND3.AMI	190 SPGKEGSHFN	200 NPYSSLFAPSI	210 ERKLIATST .:. :	220 FCWSIMLATL	230 VYLSFLVDP-' 	240 V-TVLKVYG : V: :	VPY
DESA.AMI	180	190	200	2		20	230
BNDS.AMI	250 IIFVMWLDAV	260 /TYLHHHGHDI	270 EKLPWYRGKE	280 EWSYLRGGL-	290 TTIDRDYGIFI		-
DESA.AMI	240	250	0	260	CTVHCDYPRW	280	
BND3.AMI	310 IHHLFPQIP	320 YHLVDATRA		340 REPKTSGAIP	350 THLVESLVAS	36 IKKDHYVSD	-
DESA.AMI	PHHLSVAIPS		:. : LKENWGPFLY 310		001SG 0 CHLY0 330	DPEHGYRTF(340	GSL
BND3.AMI	IVF						
DESA.AMI	KKV 350						

FIG.4



RECTIFIED SHEET (RULE 91) ISA/EP

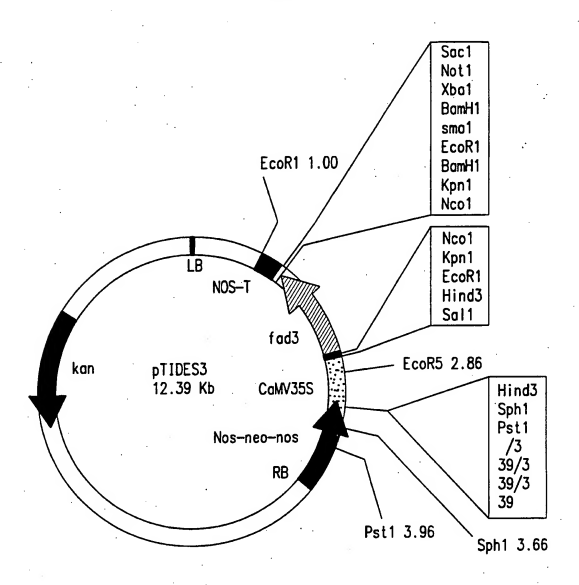


FIG.6

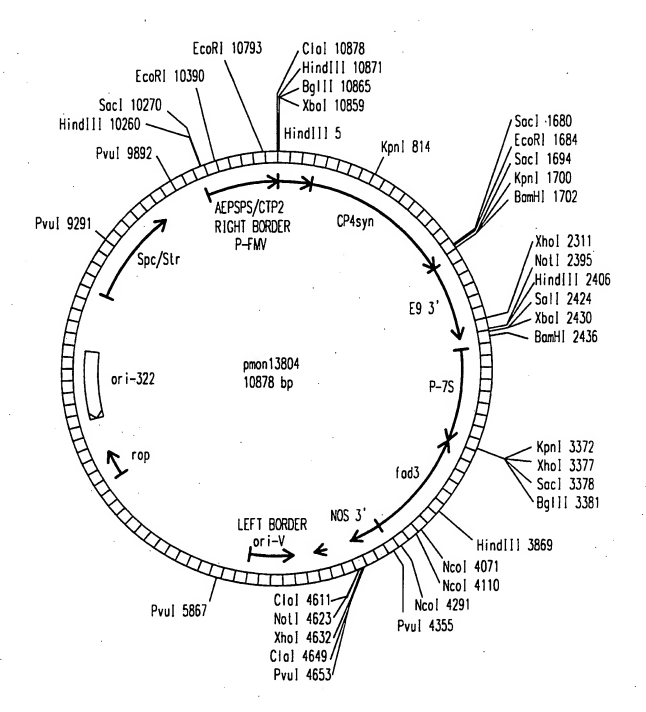


FIG.7
RECTIFIED SHEET (RULE 91)
ISA/EP

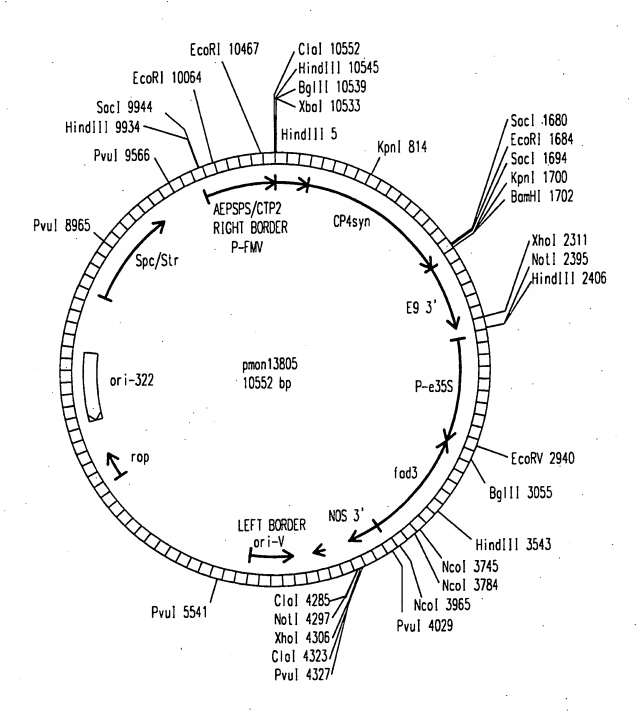
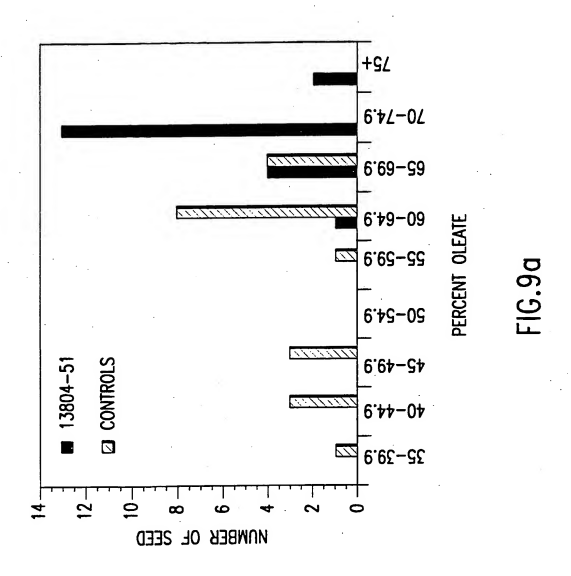
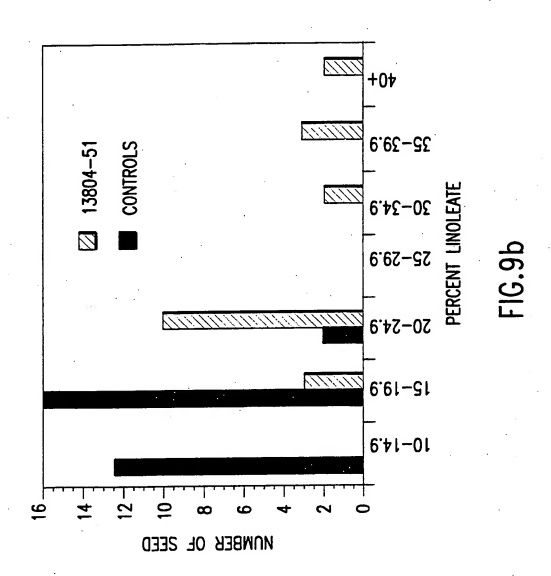


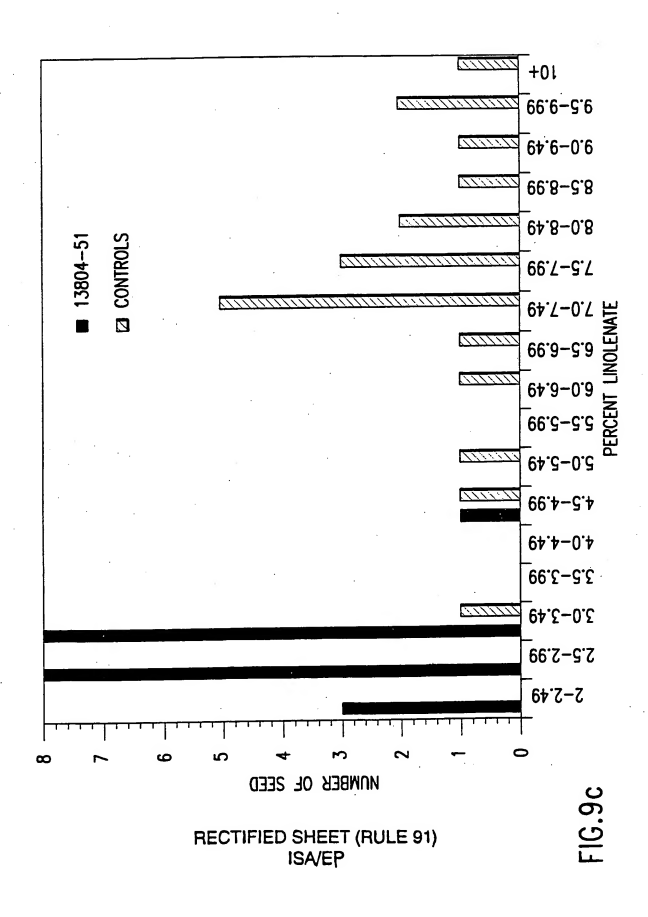
FIG.8
RECTIFIED SHEET (RULE 91)
ISA/EP



RECTIFIED SHEET (RULE 91) ISA/EP



RECTIFIED SHEET (RULE 91) ISA/EP



GGAAAACACA AGTITCTCTC ACACACATTA TCTCTTTCTC TATTACCACC ACTCATTCAT	60
AACAGAAACC CACCAAAAAA TAAAAAGAGA GACTTTTCAC TCTGGGGAGA GAGCTCAAGT	120
TCTA ATG GCG AAC TTG GTC TTA TCA GAA TGT GGT ATA CGA CCT CTC CCC Met Alo Asn Leu Vol Leu Ser Glu Cys Gly Ile Arg Pro Leu Pro 1 5 10 15	169
AGA ATC TAC ACA ACA CCC AGA TCC AAT TTC CTC TCC AAC AAC AAA Arg lie Tyr Thr Thr Pro Arg Ser Asn Phe Leu Ser Asn Asn Asn Lys 20 25 30	217
TTC AGA CCA TCA CTT TCT TCT TCT TAC AAA ACA TCA T	265
CTG TCT TTT GGT CTG AAT TCA CGA GAT GGG TTC ACG AGG AAT TGG GCG Leu Ser Phe Gly Leu Asn Ser Arg Asp Gly Phe Thr Arg Asn Trp Alo 50 55 60	313
TTG AAT GTG AGC ACA CCA TTA ACG ACA CCA ATA TTT GAG GAG TCT CCA Leu Asn Val Ser Thr Pro Leu Thr Thr Pro 11e Phe Glu Glu Ser Pro 65 70 75	361
TTG GAG GAA GAT AAT AAA CAG AGA TTC GAT CCA GGT GCG CCT CCT CCG Leu Glu Glu Asp Asn Lys Gln Arg Phe Asp Pro Gly Alo Pro Pro Pro 80 85 90 95	409
TTC AAT TTA GCT GAT ATT AGA GCA GCT ATA CCT AAG CAT TGT TGG GTT Phe Asn Leu Alo Asp lie Arg Alo Alo lie Pro Lys His Cys Trp Vol 100 105 110	457
AAG AAT CCA TGG AAG TCT TTG AGT TAT GTC GTC AGA GAC GTC GCT ATC Lys Asn Pro Trp Lys Ser Leu Ser Tyr Val Val Arg Asp Val Ala Ile 115 120 125	505
CTC TIT GCA TTG GCT GCT GGA GCT GCT TAC CTC AAC AAT TGG ATT GTT Vol Phe Alo Leu Alo Alo Gly Alo Alo Tyr Leu Asn Asn Trp lle Vol 130 135 140	553

FIG.10a

	Leu					Trp	CTC Leu	Phe	601
Leu								AAG Lys 175	649
							GTC Vol 190		697
								GGA Gly	745
							ATC lle	TAC Tyr	793
	Leu						CCT Pro		841
							GGG Gly		889
							AAA Lys 270		937
							GCT Alo		985
			Phe				CTC Leu		1033

FIG.10b

														TTT Phe		1081	
												_	_	TAC Tyr		1129).
														GAT Asp 350	-	1177	.
														CAT		1225	
														GAA Glu		1273	
													_	CCT Pro		1321	
														AAA Lys		1369	
														TAC Tyr 430		1417	
	-	_	CCA Pro 435												TGAAATGAA	NG ,	1472
CAGO	CTTG	SAG A	ATTGA	VAGT1	וו זו	TCTA	ATTIC	AG#	CCAC	CTG	ATTI	TTTC	CT 1	TACTO	STATCA	1532	
ATTI	ATTO	STG 1	CACC	CACC	CA .GA	AGAG1	TAGT	AT(стсто	SAAT	ACG/	ATCG#	ATC A	GATO	GAAAC	1592	
AACA	AAT.1	TG T	TTGC	GAT/	AC TO	SAAGO	CTATA	TA1	ACCA	AATA	AAA.		VAA A	AA		1645	

FIG.10c

Met 1	Alo	Asn	Leu	Val 5	Leu	Ser	Glu	Cys	Gly 10	He	Arg	Pro	Leu	Pro 15	Arg
lle	Tyr	Thr	Thr 20	Pro	Arg	Ser	Asn	Phe 25	Leu	Ser	Asn	Asn	Asn 30	Lys	Phe
Arg	Pro	Ser 35	Leu	Ser	Ser	Ser	Ser 40	Tyr	Lys	Thr	Ser	Ser 45	Ser	Pro	Leu
Ser	Phe 50	Gly	Leu	Asn	Ser	Arg 55	Asp	Gly	Phe	Thr	Arg 60	Asn	Trp	Alo	Leu
Asn 65	Va l	Ser	Thr	Pro	Leu 70	Thr	Thr	Pro	lle	Phe 75	Glu	Glu	Ser	Pro	Leu 80
Glu	Glu	Asp	Asn	Lys 85	GIn	Arg	Phe	Asp	Pro 90	Gly	Ala	Pro	Pro	Pro 95	Phe
Asn	Leu	Ala	Asp 100	He	Arg	Ala	Ala	I I e. 105	Pro	Lys	His	Cys	Trp 110	Val	Lys
Asn	Pro 	Trp 115	Lys	Ser	Leu	Ser	Tyr 120	Val	Val	Arg	Asp	Vol 125	AΙσ	He	Val
Phe	Ala 130	Leu	Alo	Ala	Gly	Ala 135	Ala	Туг	Leu	Asn	Asn 140	Trp	He	Val	Trp
Pro 145	Leu	Tyr	Trp	Leu	Ala 150	Gin	Gly	Thr	Met	Phe 155	Trp	Alo	Leu	Phe	Va I 160
Leu	Gly	His	Asp	Cys 165	Gly	His	Gly	Ser	Phe 170		Asn	Asp	Pro	Lys 175	Leu
Asn	Ser	Val	Va I 180	Gly	His	Leu	Leu	His 185	Ser	Ser	lle	Leu	Va l 190	Pro	Tyr
His	Gly	Trp 195	Arg	He	Ser	His	Arg 200	Thr	His	His	GIn	Asn 205	His	Gly	His
Val	Glu 210	Asn	Asp	Glu	Ser	Trp 215	His	Pro	Met	Ser	Glu 220	Lys	He.	Tyr	Asn

FIG. 11a RECTIFIED SHEET (RULE 91) ISA/EP

Thr 225	Leu	Asp	Lys	Pro	Thr 230	Arg	Phe	Phe	Arg	Phe 235		Leu	Pro	Leu	Va I 240
Met	Leu	Ala	Tyr	Pro 245	Phe	Tyr	Leu	Trp	A1 o 250	Arg	Ser	Pro	Gly	Lys 255	Lys
Gly	Ser	His	Tyr 260	His	Pro	Asp	Ser	Asp 265	Leu	Phe	Leu	Pro	Lys 270	Glu	Arg
Lys	Asp	Vol 275	Leu	Thr	Ser	Thr	A1 a 280	Cys	Trp	Thr	Alo	Me t 285	Alo	Ala	Leu
Leu	Va I 290	Cys	Leu	Asn	Phe	Thr 295	He	Gly	Pro	He	GIn 300	Met	Leu	Lys	Leu
Tyr 305	Gly	He	Pro	Tyr	Trp 310	lle	Asn	Val	Met	Trp 315	Leu	Asp	Phe	Val	Thr 320
Tyr	Leu	His	His	His 325	-	His	Glu	Asp	Lys 330	Leu	Pro	Trp	Tyr	Arg 335	Gly
Lys	Glu	Trp	Ser 340	Tyr	Leu	Arg	Gly	G I y 345	Leu	Thr	Thr	Leu	Asp 350	Arg	Asp
Tyr	Gly	Leu 355	lie	Asn	Asn	lle	His 360	His	Asp	He	Gly	Thr 36 5	His	Val	lie
His	His 370	Leu	Phe	Pro	Gln	11e 375	Pro	His	Tyr	His	Leu 380	Val	Glu	Alo	Thr
Glu 385	Alo	Alo	Lys	Pro	Va I 390		Gly	Lys	Tyr	Tyr 395		Glu	Pro		Lys 400
Ser	Gly	Pro	Leu	Pro 405	Leu	His	Leu	Leu	Glu 410	He	Leu	Ala	Lys	Ser 415	He
Lys	Glu	Asp	His 420	Туг	Va I	Ser	Asp	Glu 425	Gly	Glu	Val	Val	Tyr 430	Tyr	Lys
Ala	Asp	Pro 435	Asn	Leu	Tyr	Gly	G I u 440	Val	Lys	Val	Arg	A1 o 445	Asp	•	

FIG.11b

AGA	GAGT	GCA /	AATA(GAAC	GA C	AGAG	ACTT	1 11	CCTC	וווד יי	CTT	CTTG	GGA A	AGAG	GCTCCA		60
			TCG Ser														108
			AAA Lys 20													1	156
		_	AAT Asn													2	204
			TTC Phe													2	252
			ACA Thr														300
			GGT													3	348
			AAG Lys 100													3	96
			AGA Arg													4	44
			AAC Asn													4	92
-			TTC Phe													.5	40

FIG.12a

GGT Gly	AGC Ser	TTC Phe	TCG Ser	AAT Asn 165	GAT Asp	CCG Pro	AGG Arg	CTG Leu	AAC Asn 170	AGT Ser	GTG Val	GCT Alo	GGT Gly	CAT His 175	CTT Leu	588	
CTT Leu	CAT His	TCC Ser	TCA Ser 180	ATT 11e	CTG Leu	GTC Val	CCT Pro	TAC Tyr 185	CAT His	GGC Gly	TGG Trp	AGG Arg	ATT He 190	AGC Ser	CAC	636	
AGA Arg	ACT	CAC His 195	CAC His	CAG GIn	AAC Asn	CAT His	GGT Gly 200	CAT His	GTC Val	GAG Glu	AAT Asn	GAC Asp 205	GAA Glu	TCA Ser	TGG Trp	684	
				GAA Glu												732	
ATG Met 225	TTT Phe	AGG Arg	TTT Phe	ACA Thr	CTG Leu 230	CCT Pro	TTT Phe	CCA Pro	ATG Met	CTC Leu 235	GCA Ala	TAC Tyr	CCT Pro	TTC Phe	TAC Tyr 240	780	
TTG Leu	TGG Trp	AAC Asn	AGA Arg	AGT Ser 245	CCA Pro	GGG Gly	AAA Lys	CAA GIn	GGT GTy 250	TCT Ser	CAT	TAT Tyr	CAT His	CCG Pro 255	GAC Asp	828	
				CTT Leu												876	
GCC Alo	TGT Cys	TGG Trp 275	ACT Thr	GCA Ala	ATG Met	GCT Alo	GCT Alo 280	TTG Leu	CTT Leu	GTT Vol	TGT Cys	CTC Leu 285	AAC Asn	TTT Phe	GTC Val	924	
				CAG Gin											ATA 11e	972	
TTT Phe 305	Val	ATG Met	TGG Trp	IIG Leu	GAC Asp 310	Phe	GTC Val	ACT Thr	TAC Tyr	TTG Leu 315	His	CAC His	CAT His	GGA Gly	CAT His 320	1020	

FIG. 12b RECTIFIED SHEET (RULE 91) ISA/EP

								GGA Gly								1068
								GAC Asp 345								1116
								ATA 11e								1164
								ACA Thr								1212
								AAC Asn								1260
								ATG Met							AGC Ser	1308
								GAG Glu 425								1356
	AGA Arg		TGA	GGAC	ATA (CTGC/	AGTG/	AA C(CAGG	CAGA(C AAC	STTA(CATA			1405
AAT	TCAT	CTT (GGCC	CATTO	CA T	TATG	TTCT	TT.	IGTT	TTGG	TGT	AAAG(CCT .	TTTC	GAGATT	1465
AAA	AAAG(CAT	Taat	TTGT	AG A	AACC	rgtg(G TA	AAAC1	TCTC	GAT(CAAA	IGA /	AATA/	CATAT	1525

FIG.12c

Met 1	Ala	Ser	Ser	Val 5	Leu	Ser	Glu	Cys	Gly 10	Phe	Arg	Pro	Leu	Pro 15	Arg
Phe	Tyr	Pro	Lys 20	His	Thr	Thr	Ser	Phe 25	Ala	Ser	Asn	Pro	Lys 30	Pro	Thr
Phe	Lys	Phe 35	Asn	Pro	Pro	Leu	Lys 40	Pro	Pro	Ser	Ser	Leu 45	Leu	Asn	Ser
Arg	Tyr 50	Gly	Phe	Tyr	Ser	Lys 55	Thr	Arg	Asn	Trp	A1 a 60	Leu	Asn	Val	Ala
Thr 65	Pro	Leu	Thr	Thr	Leu 70	Gln	Ser	Pro	Ser	G1u 75	Glu	Asp	Thr	Glu	Arg 80
Phe	Asp	Pro	Gly	A1 a 85	Pro	Pro	Pro	Phe	Asn 90	Leu	Ala	Asp	lle	Arg 95	Alo
Ala	He	Pro	Lys 100	His	Cys	Trp	Val	Lys 105	Asn	Pro ·	Trp	Met	Ser 110	Met	Ser
Tyr	Vol	Va I 115	Arg	Asp	Val	Ala	lie 120	Val	Phe	Gly	Leu	Ala 125	Ala	Vol	Ala
Ala	Tyr 130	Phe	Asn	Asn	Тгр	Leu 135	Leu	Trp	Pro	Leu	Tyr 140	Trp	Phe	Ala	Gin
G1y 145	Thr	Met	Phe	Trp	Alo 150	Leu	Phe	Val	Leu	G1y 155	His	Asp	Cys	Gly	His 160
Gly	Ser	Phe	Ser	Asn 165	Asp	Pro	Arg	Leu	Asn 170	Ser	Val	Ala	Gly	His 175	Leu
Leu	His	Ser	Ser 180	He	Leu	Val	Pro	Tyr 185	His	Gly	Trp	Arg	11e 190	Ser	His
Arg	Thr	His 195	His	GIn	Asn	His	Gly 200	His	Val	Glu	Asn	Asp 205	Glu	Ser	Trp
His	Pro 210	Leu	Pro	Glu	Ser	lle 215	Tyr	Lys	Asn	Leu	Glu 220	Lys	Thr	Thr	Gin

Met 225	Phe	Arg	Phe	Thr	Leu 230	Pro	Phe	Pro	Met	Leu 235	Alo	Tyr	Pro	Phe	Tyr 240
Leu	Trp	Asn	Arg	Ser 245	Pro	Gly	Lys	GIn	Gly 250	Ser	His	Tyr	His	Pro 255	Asp
Ser	Asp	Leu	Phe 260	Leu	Pro	Lys	Glu	Lys 265	Ļys	Asp	Val	Leu	Thr 270	Ser	Thr
Ala	Cys	Trp 275	Thr	Ala	Met	Αlσ	Ala 280	Leu	Leu	Val	Cys	Leu 285	Asn	Phe	Val
Met	Gly 290	Pro	lie	Gļn	Met	Leu 295	Lys	Leu	Tyr	Gly	11e 300	Pro	Tyr	Trp	He
Phe 305	Vol	Met	Trp	Leu	Asp 310	Phe	Vol	Thr	Tyr	Leu 315		His	His	Gly	His 320
Glu	Asp	Lys	Leu	Pro 32 5	Trp	Tyr	Arg	Gly	Lys 330	Glu	Trp	Ser	Tyr	Leu 335	Arg
Gly	Gly	Leu	Thr 340	Thr	Leu	Asp		Asp .345	Tyr	Gly	Trp	lle	Asn 350	Asn	Пe
His	His	Asp 355	lle	Gly	Thr	His	Va I 360	He	His	His	Leu	Phe 365	Pro	GIn	He
Pro	His 370	Tyr	His	Leu	Vol	Glu 375		Thr	Glu	Ala	A1 a 380	Lys	Pro	Val	Leu
G1y 385	Lys	Tyr	Tyr	Arg	Glú 390	Pro	Lys	Asn	Ser	Gly 395	Pro	Leu	Pro	Leu	His 400
Leu	Leu	Gly	Ser	Leu 405		Lys	Ser	Met	Lys 410	GIn	Asp	His	Phe	Val 415	Ser
Asp	Thr	Gly	Asp 420		Val	Tyr	Tyr	G1u 425		Asp	Pro	Lys	Leu 430	Asn	Gly
Gin	Arg	Thr													

Inte. 1al Application No PCT/US 94/01321

A. CLASSIFICATION OF SUBJECT MATTER IPC 5 C12N15/82 C12N1 C12N15/11 C12N5/10 A01H5/00 C12N15/53 C11B1/00 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 5 C12N A01H C11B Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Citation of document, with indication, where appropriate, of the relevant passages Category * 37 X SCIENCE vol. 258 , 20 November 1992 , LANCASTER, pages 1353 - 1355 ARONDEL, V., ET AL. 'Map-based cloning of a gene controlling omega-3 fatty acid desaturation in Arabidopsis' 1,2,4,8, Y see the whole document 9,11,17, 18,22, 23,25, 29,30, 32,38 Patent family members are listed in annex. Further documents are listed in the continuation of box C. Special categories of cited documents: "I" later document published after the international filing date or priority date and not in conflict with the application but used to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to filing date involve an inventive step when the document is taken alone 'L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another "Y" document of particular relevance; the claimed invention citation or other special reason (as specified) cannot be considered to involve an inventive step when the document is combined with one or more other such docudocument referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled other means document published prior to the international filing date but "&" document member of the same patent family later than the priority date claimed Date of mailing of the international search report Date of the actual completion of the international search 1 4 -06- 1994 1 June 1994 Authorized officer Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Ripswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax (+31-70) 340-3016 Maddox, A

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